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### **PCT**

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#### INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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(54) Title: PROSTATE-SPECIFIC MEMBRANE ANTIGEN AND USES THEREOF

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#### (57) Abstract

This invention provides an isolated mammalian nucleic acid molecule encoding an alternatively spliced prostate-specific membrane (PSM') antigen. This invention provides an isolated nucleic acid molecule encoding a prostate-specific membrane antigen promoter. This invention provides a method of detecting hematogenous micrometastic tumor cells of a subject, and determining prostate cancer progression in a subject.

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#### PROSTATE-SPECIFIC MEMBRANE ANTIGEN AND USES THEREOF

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This application is a continuation-in-part of United States Application Serial Nos. 08/466,381 and 08/470,735, both filed June 2, 1995, which are continuations of U.S. Serial No. 08/394,152, filed February 24, 1995, the contents of which are hereby incorporated by reference.

This invention disclosed herein was made in part with Government support under NIH Grants No. DK47650 and CA58192, CA-39203, CA-29502, CA-08748-29 from the Department of Health and Human Services. Accordingly, the U.S. Government has certain rights in this invention.

#### 20 BACKGROUND OF THE INVENTION

Throughout this application various references are referred to within parentheses. Disclosures of these publications in their entireties are hereby incorporated by reference into this application to more fully describe the state of the art to which this invention pertains. Full bibliographic citation for these references may be found at the end of each set of Examples in the Experimental Details section.

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Prostate cancer is among the most significant medical problems in the United States, as the disease is now the most common malignancy diagnosed in American males. In 1992 there were over 132,000 new cases of prostate cancer detected with over 36,000 deaths attributable to the disease, representing a 17.3% increase over 4 years (2). Five year survival rates for patients with prostate cancer range from 88% for those with localized disease to 29% for those with metastatic disease. The

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rapid increase in the number of cases appears to result in part from an increase in disease awareness as well as the widespread use of clinical markers such as the secreted proteins prostate-specific antigen (PSA) and prostatic acid phosphatase (PAP) (37).

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The prostate gland is a site of significant pathology affected by conditions such as benign growth (BPH), cancer) and infection neoplasia (prostatic (prostatitis). Prostate cancer represents the second leading cause of death from cancer in man (1). However prostatic cancer is the leading site for cancer development in men. The difference between these two facts relates to prostatic cancer occurring with increasing frequency as men age, especially in the ages beyond 60 at a time when death from other factors often spectrum of biologic Also. the intervenes. aggressiveness of prostatic cancer is great, so that in some men following detection the tumor remains a latent histologic tumor and does not become clinically significant, whereas in other it progresses rapidly, metastasizes and kills the man in a relatively short 2-5 year period (1, 3).

In prostate cancer cells, two specific proteins that 25 are made in very high concentrations are prostatic acid phosphatase (PAP) and prostate specific antigen (PSA) (4, 5, 6). These proteins have been characterized and have been used to follow response to therapy. With the development of cancer, the normal architecture of the 30 gland becomes altered, including loss of the normal duct structure for the removal of secretions and thus the secretions reach the serum. Indeed measurement of serum PSA is suggested as a potential screening method for prostatic cancer. Indeed, the relative amount of 35 PSA and/or PAP in the cancer reduces as compared to normal or benign tissue.

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PAP was one of the earliest serum markers for detecting metastatic spread (4). PAP hydrolyses tyrosine phosphate and has a broad substrate specificity. Tyrosine phosphorylation is often increased with oncogenic transformation. It has been hypothesized that during neoplastic transformation there is less phosphatase activity available to inactivate proteins that are activated by phosphorylation on tyrosine residues. In some instances, insertion of phosphatases that have tyrosine phosphatase activity has reversed the malignant phenotype.

PSA is a protease and it is not readily appreciated how loss of its activity correlates with cancer development 15 (5, 6). The proteolytic activity of PSA is inhibited by zinc. Zinc concentrations are high in the normal prostate and reduced in prostatic cancer. Possibly the loss of zinc allows for increased proteolytic activity As proteases are involved in metastasis and 20 some proteases stimulate mitotic activity, potentially increased activity of PSA could be hypothesized to play a role in the tumors metastases and spread (7).

- Both PSA and PAP are found in prostatic secretions.

  Both appear to be dependent on the presence of androgens for their production and are substantially reduced following androgen deprivation.
- Prostate-specific membrane antigen (PSM) which appears to be localized to the prostatic membrane has been identified. This antigen was identified as the result of generating monoclonal antibodies to a prostatic cancer cell, LNCaP (8).

Dr. Horoszewicz established a cell line designated LNCaP from the lymph node of a hormone refractory,

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heavily pretreated patient (9). This line was found to have an aneuploid human male karyotype. It maintained prostatic differentiation functionality in that it produced both PSA and PAP. It possessed an androgen receptor of high affinity and specificity. Mice were immunized with LNCaP cells and hybridomas were derived from sensitized animals. A monoclonal antibody was derived and was designated 7E11-C5 (8). The antibody staining was consistent with a membrane location and isolated fractions of LNCaP cell membranes exhibited a strongly positive reaction with immunoblotting and This antibody did not inhibit or ELISA techniques. enhance the growth of LNCaP cells in vitro or in vivo. The antibody to this antigen was remarkably specific to prostatic epithelial cells, as no reactivity was observed in any other component. Immunohistochemical staining of cancerous epithelial cells was more intense than that of normal or benign epithelial cells.

Dr. Horoszewicz also reported detection 20 immunoreactive material using 7E11-C5 in serum of prostatic cancer patients (8). The immunoreactivity was detectable in nearly 60% of patients with stage D-2 disease and in a slightly lower percentage of patients with earlier stage disease, but the numbers of patients 25 in the latter group are small. Patients with benign prostatic hyperplasia (BPH) were negative. with no apparent disease were negative, but 50-60% of patients in remission yet with active stable disease or demonstrated positive 30 with progression serum reactivity. Patients with non prostatic tumors did not show immunoreactivity with 7E11-C5.

The 7E11-C5 monoclonal antibody is currently in clinical trials. The aldehyde groups of the antibody were oxidized and the linker-chelator glycol-tyrosyl(n, \epsilon-diethylenetriamine-pentacetic acid)-lysine (GYK-

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DTPA) was coupled to the reactive aldehydes of the heavy chain (10). The resulting antibody was designated CYT-356. Immunohistochemical staining patterns were similar except that the CYT-356 modified antibody stained skeletal muscle. The comparison of CYT-356 with 7E11-C5 monoclonal antibody suggested both had binding to type 2 muscle fibers. The reason for the discrepancy with the earlier study, which reported skeletal muscle to be negative, was suggested to be due to differences in tissue fixation techniques. Still, the most intense and definite reaction was observed with prostatic epithelial cells, especially cancerous Reactivity with mouse skeletal muscle was detected with immunohistochemistry but not in imaging The Indium 111-labeled antibody localized to LNCaP tumors grown in nude mice with an uptake of nearly 30% of the injected dose per gram tumor at four In-vivo, no selective retention of the antibody was observed in antigen negative tumors such as PC-3 and DU-145, or by skeletal muscle. Very little was known about the PSM antigen. An effort at purification and characterization has been described at meetings by Dr. George Wright and colleagues (11, 12).

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#### BRIEF DESCRIPTION OF THE FIGURES

Figure 1: Signal in lane 2 represent the 100kD PSM antigen. The EGFr was used as the positive control and is shown in lane 1. Incubation with rabbit antimouse (RAM) antibody alone served as negative control and is shown in lane 3.

10 Figures 2A-2D: Upper two photos show LNCaP cytospins staining positively for PSM antigen.

Lower left in DU-145 and lower right is PC-3 cytospin, both negative for PSM antigen expression.

Figures 3A-3D: Upper two panels are human prostate sections (BPH) staining positively for PSM antigen. The lower two panels show invasive prostate carcinoma human sections staining positively for expression of the PSM antigen.

Figure 4: 100kD PSM antigen following immunoprecipitation of <sup>35</sup>S-Methionine labelled LNCaP cells with Cyt-356 antibody.

Figure 5: 3% agarose gels stained with Ethidium bromide revealing PCR products obtained using the degenerate PSM antigen primers. The arrow points to sample IN-20, which is a 1.1 kb PCR product which was later confirmed to be a partial cDNA coding for the PSM gene.

Figures 6A-6B: 2% agarose gels of plasmid DNA

resulting from TA cloning of PCR products. Inserts are excised from the PCR II vector (Invitrogen Corp.) by digestion with EcoRI. 1.1 kb PSM gene partial cDNA product is shown in lane 3 of gel 1.

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Autoradiogram showing size of cDNA represented in applicants' LNCaP library using M-MLV reverse transcriptase.

Figure 8:

Figure 7:

Restriction analysis of full-length clones of PSM gene obtained after screening cDNA library. Samples have been cut with Not I and Sal I restriction enzymes to liberate the insert.

20 Figure 9:

Plasmid Southern autoradiogram of full length PSM gene clones. Size is approximately 2.7 kb.

Figure 10:

Northern blot revealing PSM expression limited to LNCaP prostate cancer line and H26 Ras-transfected LNCaP cell line. PC-3, DU-145, T-24, SKRC-27, HELA, MCF-7, HL-60, and others were are all negative.

(kb), and 28S and 18S ribosomal RNA

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Figure 11: Autoradiogram of Northern analysis revealing expression of 2.8 kb PSM message unique to the LNCaP cell line (lane 1), and absent from the DU-145 (lane 2) and PC-3 cell lines (lane 3). RNA size ladder is shown on the left

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bands are indicated on the right.

#### Figures 12A-12B:

Results of PCR of human prostate tissues using PSM gene primers. Lanes are numbered from left to right. Lane 1, LNCaP; Lane 2, H26; Lane 3, DU-145; Lane 4, Normal Prostate; Lane 5, BPH; Lane 6, Prostate Cancer; Lane 7, BPH; Lane 8, Normal; Lane 9, BPH; Lane 10, BPH; Lane 11, BPH; Lane 12, Normal; Lane 13, Normal; Lane 14, Cancer; Lane 15, Cancer; Lane 16, Cancer; Lane 17, Normal; Lane 18, Cancer; Lane 19, IN-20 Control; Lane 20, PSM cDNA

Figure 13: Isoelectric point of PSM antigen (non-glycosylated)

20 Figures 14:1-8 Secondary structure of PSM antigen

#### Figures 15A-15B:

A. Hydrophilicity plot of PSM antigen
B. Prediction of membrane spanning
segments

#### Figures 16:1-11

Homology with chicken, rat and human transferrin receptor sequence.

Figures 17A-17C:

Immunohistochemical detection of PSM antigen expression in prostate cell lines. Top panel reveals uniformly high level of expression in LNCaP cells; middle panel and lower panel are DU-145 and PC-3 cells respectively,

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both negative.

Figure 18: Autoradiogram of protein gel revealing products of PSM coupled in-vitro transcription/translation. Non-glycosylated PSM polypeptide is seen at 84 kDa (lane 1) and PSM glycoprotein

synthesized following the addition of microsomes is seen at 100 kDa (lane 2).

Figure 19:

Western Blot analysis detecting PSM in expression transfected non-PSM expressing PC-3 cells. 100 kDa PSM glycoprotein species is clearly seen in LNCaP membranes (lane 1), LNCaP crude lysate (lane 2), and PSM-transfected PC-3 cells (lane 4), but undetectable in native PC-3 cells (lane 3).

human brain (lane 4) and human salivary

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Pigure 20: Autoradiogram of ribonuclease protection gel assaying for PSM mRNA expression in normal human tissues. Radiolabeled 1 kb DNA ladder (Gibco-BRL) is shown in lane 1. Undigested probe is 400 nucleotides (lane 2), expected protected PSM band is 350 nucleotides, and tRNA control is shown (lane 3). A strong signal is seen in human prostate (lane 11), with very faint, but detectable signals seen in

35 **Figure 21:** Autoradiogram of ribonuclease protection gel assaying for PSM mRNA expression in LNCaP tumors grown in

gland (lane 12).

nude mice, and in human prostatic tissues. 32P-labeled 1 kb DNA ladder is shown in lane 1. 298 nucleotide undigested probe is shown (lane 2), and tRNA control is shown (lane 3). mRNA expression is clearly detectable in LNCaP cells (lane 4), orthotopically grown LNCaP tumors in nude mice with and without matrigel (lanes 5 and 6), and subcutaneously implanted and grown LNCaP tumors in nude mice (lane 7). PSM mRNA expression is also seen in normal human prostate (lane 8), and in moderately differentiated human prostatic adenocarcinoma (lane 10). Very faint expression is seen in a sample of human prostate tissue with benign hyperplasia (lane 9).

20 Figure 22:

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Ribonuclease protection assay for PSM expression in LNCaP cells treated with physiologic doses of various steroids for 24 hours. 32P-labeled DNA ladder is shown in lane 1. 298 nucleotide undigested probe is shown (lane 2), and tRNA control is shown (lane 3). mRNA expression is highest in untreated LNCaP cells in charcoal-stripped media (lane 4). Applicant see significantly diminished PSM expression in LNCaP cells treated with DHT (lane Testosterone (lane 6), Estradiol (lane 7), and Progesterone (lane 8), with little response to Dexamethasone (lane 9).

Figure 23: Data illustrating results of PSM DNA

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and RNA presence in transfect Dunning cell lines employing Southern and Northern blotting techniques

#### 5 Figures 24A-24B:

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Figure A indicates the power of cytokine transfected cells to teach unmodified cells. Administration was directed to the parental flank or prostate cells. The results indicate the microenvironment considerations.

Figure B indicates actual potency at a particular site. The tumor was implanted in prostate cells and treated with immune cells at two different sites.

#### Figures 25A-25B:

20 Relates potency of cytokines in inhibiting growth of primary tumors. Animals administered un-modified parental tumor cells and administered vaccine transfected 25 Following prostatectomy of rodent tumor results in survival increase.

Figure 26: PCR amplification with nested primers improved the level of detection of prostatic cells from approximately one prostatic cell per 10,000 MCF-7 cells to better than one cell per million MCF-7 cells, using either PSA.

35 Figure 27: PCR amplification with nested primers improved the level of detection of prostatic cells from approximately one

prostatic cell per 10,000 MCF-7 cells to better than one cell per million MCF-7 cells, using PSM-derived primers.

- 5 Figure 28:
- A representative ethidium stained gel photograph for PSM-PCR. Samples run in lane A represent PCR products generated from the outer primers and samples in lanes labeled B are products of inner primer pairs.

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Figure 29:

PSM Southern blot autoradiograph. The sensitivity of the Southern blot analysis exceeded that of ethidium staining, as can be seen in several samples where the outer product is not visible on figure 3, but is detectable by Southern blotting as shown in figure 4.

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Figure 30: Characteristics of the 16 patients analyzed with respect to their clinical stage, treatment, serum PSA and PAP values, and results of assay.

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#### Figures 31A-31D:

The DNA sequence of the 3 kb XhoI fragment of p683 which includes 500 bp of DNA from the RNA start site was determined Sequence 683XFRVS starts from the 5' distal end of PSM promoter.

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Figure 32: Potential binding sites on the PSM promoter.

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Figure 33: Promoter activity of PSM up-stream fragment/CAT gene chimera.

Figure 34:

Comparison between PSM and PSM' cDNA. Sequence of the 5' end of PSM cDNA (5) is shown. Underlined region denotes nucleotides which are present in PSM cDNA sequence but absent in PSM' cDNA. Boxed region represents the putative transmembrane domain of PSM antigen.

\* Asterisk denotes the putative translation initiation site for PSM'.

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Figure 35:

Graphical representation of PSM and PSM' cDNA sequences and antisense PSM RNA probe (b). PSM cDNA sequence with complete coding region (5). (a) PSM' cDNA sequence from this study. (c) Cross hatched and open boxes denote sequences identity in PSM and PSM'. Hatched box indicates sequence absent from PSM'. Regions of cDNA sequence complementary to the antisense probe are indicated by dashed lines between the sequences.

Figure 36:

RNase protection assay with **PSM** in primary prostatic specific probe tissues. Total cellular RNA was isolated from human prostatic samples: normal prostate, BPH, and CaP. PSM and spliced variants are indicated with arrows at right. The left lane is a DNA ladder. Samples from different patients are classified as: lanes 3-6, CaP, carcinoma of prostate; BPH, benign prostatic hypertrophy, lanes normal, normal prostatic tissue, lanes Autoradiograph was exposed for longer period to read lanes 5 and 9.

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5	Figure 37:	Tumor Index, a quantification of the expression of PSM and PSM'. Expression of PSM and PSM' (Fig.3) was quantified by densitometry and expressed as a ratio of PSM/PSM' on the Y-axis. Three samples each were quantitated for					
10		primary CaP, BPH and normal prostate tissues. Two samples were quantitated for LNCaP. <i>Normal</i> , normal prostate tissue.					
	Figure 38:	Characterization of PSM membrane bound and PSM' in the cytosol.					

15 Figure 39: Intron 1F: Forward Sequence. Intron 1 contains a number of trinucleotide repeats which can be area associated with chromosomal instability in tumor cells. LNCaP cells and primary prostate tissue are identical, however in the PC-3 and Du-145 tumors they have substantially altered levels of these trinucleotide repeats which may relate to their lack of expression of PSM.

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#### Figures 40A-40B:

Intron 1R: Reverse Sequence

Figure 41: Intron 2F: Forward Sequence

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Figure 42: Intron 2R: Reverse Sequence

Figures 43A-43B:

Intron 3F: Forward Sequence

35 Figures 44A-44B:

Intron 3R: Reverse Sequence

Figures 45A-45B:

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Intron 4F: Forward Sequence

#### Figures 46A-46B:

Intron 4R: Reverse Sequence

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#### Figures 47A-47D:

Sequence of the genomic region upstream of the 5' transcription start site of PSM.

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Figure 48:

Photograph of ethidium bromide stained gel depicting representative negative and positive controls used in the Samples 1-5 were study. respectively: male with prostatis, a healthy female volunteer, a male with BPH, a control 1:1,000,000 dilution of LNCaP cells, and a patient with renal cell carcinoma. Below each reaction is control reaction the corresponding with beta-2-microglobulin performed primers to assure RNA integrity. PCR products were detected for any of these negative controls.

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Figure 49:

Photograph of gel displaying representative positive PCR results using PSM primers in selected patients with either localized or disseminated prostate cancer. Sample 1-5 were from. respectively: a patient with clinically localized stage Tl, disease, a radical patient with prostatectomy confined disease and a negative serum PSA, a radical prostatectomy patient with locally advanced disease and a negative serum PSA, a patient with

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treated stage D2 disease, and a patient with treated hormone refractory disease.

5 **Figure 50:** Chromosomal location of PSM based on cosmid construction.

Figure 51:

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Human monochromosomal somatic cell hybrid blot showing that chromosome 11 contained the PSM genetic sequence by Southern analysis. DNA panel digested with PstI restriction enzyme and probed with PSM cDNA. Lanes M and H refer to mouse and hamster DNAs. The numbers correspond to the human chromosomal DNA in that hybrid.

Figure 52: Ribonuclease protection assay using PSM radiolabeled RNA probe revels an abundant PSM mRNA expression in AT6.1-11 clone 1, but not in AT6.1-11 clone 2, thereby mapping PSM to 11p11.2-13 region.

25 **Figure 53:** Tissue specific expression of PSM RNA by Northern blotting and RNAse protection assay.

Figure 54: Mapping of the PSM gene to the llpll.2pl3 region of human chromosome 11 by
southern blotting and in-situ
hybridization.

Figure 55: Schematic of potential response elements.

Figure 56: Genomic organization of PSM gene.

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Figure 57: Schematic of metastatic prostate cell

#### Figure 58A-58C:

Nucleic acid of PSM genomic DNA is read

5 prime away from the transcription
start site: number on the sequences
indicates nucleotide upstream from the
start site. Therefore, nucleotide #121

10 is actually -121 using conventional
numbering system.

#### Figure 59:

Representation of NAAG 1, acividin, azotomycin, and 6-diazo-5-oxo-norleucine, DON.

#### Figure 60:

Preparation of N-acetylaspartylglutamate, NAAG 1.

#### Figure 61:

Synthesis of N-acetylaspartylglutamate, NAAG 1.

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Figure 62:

Synthesis of N-phosphonoacetylaspartyl-L-glutamate.

30 Figure 63:

Synthesis of 5-diethylphosphonon-2 amino benzylvalerate intermediate.

Figure 64:

35 Synthesis of analog 4 and 5.

Figure 65:

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Representation of DON, analogs 17-20.

5 Figure 66:

Substrates for targeted drug delivery,

analog 21 and 22.

Figure 67:

Dynemycin A and its mode of action.

Figure 68:

Synthesis of analog 28.

15 Figure 69:

Synthesis for intermediate analog 28.

Figure 70:

Attachment points for PALA.

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Figure 71:

Mode of action for substrate 21.

Figures 72A-72D:

25 Intron 1F: Forward Sequence.

Figures 73A-73E:

Intron 1R: Reverse Sequence

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Intron 2F: Forward Sequence

Figures 75A-75C:

Intron 2R: Reverse Sequence

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Figures 76A-76B:

Intron 3F: Forward Sequence

Figures 77A-77B:

Intron 3R: Reverse Sequence

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Intron 4F: Forward Sequence

Figures 79A-79E:

Intron 4RF: Reverse Sequence

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Figure 80:

PSM genomic organization of the exons and 19 intron junction sequences. The exon/intron junctions (See Example 15) are as follows:

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- Exon /intron 1 at bp 389-390;
- Exon /intron 2 at bp 490-491;
- 3. Exon /intron 3 at bp 681-682;
- Exon /intron 4 at bp 784-785;
- 5. Exon /intron 5 at bp 911-912;
- Exon /intron 6 at bp 1096-1097;
- Exon /intron 7 at bp 1190-1191;
- Exon /intron 8 at bp 1289- 1290;
- 9. Exon /intron 9 at bp 1375-1376;
- 25 10. Exon /intron 10 at bp 1496-1497;
  - 11. Exon /intron 11 at bp 1579-1580;
  - 12. Exon /intron 12 at bp 1640-1641;
  - 13. Exon /intron 13 at bp 1708-1709;
  - 13. Exon /Intron 13 at bp 1700 1703;
    14. Exon /intron 14 at bp 1803-1804;
  - 15. Exon /intron 15 at bp 1892-1893;
  - 15. DAON / INCION 15 do ap acces
  - Exon /intron 16 at bp 2158-2159;
  - 17. Exon /intron 17 at bp 2240-2241;
  - 18. Exon /intron 18 at bp 2334-2335;
  - 19. Exon /intron 19 at bp 2644-2645.

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DNA molecule of an isolated mammalian nucleic acid molecule encoding a mammalian prostate-specific membrane antigen. This invention also provides an isolated mammalian cDNA molecule encoding a mammalian prostate-specific membrane antigen. This invention provides an isolated mammalian RNA molecule encoding a mammalian prostate-specific membrane antigen.

In the preferred embodiment of this invention, the isolated nucleic sequence is cDNA from human as shown in Figures 47A-47D. This human sequence was submitted to GenBank (Los Alamos National Laboratory, Los Alamos, New Mexico) with Accession Number, M99487 and the description as PSM, Homo sapiens, 2653 base-pairs.

This invention also encompasses DNAs and cDNAs which encode amino acid sequences which differ from those of PSM or PSM' antigen, but which should not produce phenotypic changes. Alternatively, this invention also encompasses DNAs and cDNAs which hybridize to the DNA and cDNA of the subject invention. Hybridization methods are well known to those of skill in the art.

For example, high stringent hybridization conditions are selected at about 5° C lower than the thermal melting point (Tm) for the specific sequence at a The Tm is the defined ionic strength and pH. temperature (under defined ionic strength and pH) at which 50% of the target sequence hybridizes to a stringent Typically, probe. matched perfectly the salt which in those be will conditions concentration is at least about 0.02 molar at pH 7 and the temperature is at least about 60°C. factors may significantly affect the stringency of others, among including, hybridization, composition and size of the complementary strands, the presence of organic solvents, ie. salt or formamide

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concentration, and the extent of base mismatching, the combination of parameters is more important than the absolute measure of any one. For Example high stringency may be attained for example by overnight hybridization at about 68°C in a 6x SSC solution, washing at room temperature with 6x SSC solution, followed by washing at about 68°C in a 6x SSC in a 0.6x SSX solution.

Hybridization with moderate stringency may be attained 10 1) filter pre-hybridizing and for example by: hybridizing with a solution of 3x sodium chloride, sodium citrate (SSC), 50% formamide, 0.1M Tris buffer 7.5, 5x Denhardt's solution; hybridization at 37°C for 4 hours; 3) hybridization at 15 37°C with amount of labelled probe equal to 3,000,000 cpm total for 16 hours; 4) wash in 2x SSC and 0.1% SDS 5) wash 4x for 1 minute each at room solution: temperature at 4x at 60°C for 30 minutes each; and 6) dry and expose to film. 20

> The DNA molecules described and claimed herein are information which they provide for the useful concerning the amino acid sequence of the polypeptide and as products for the large scale synthesis of the polypeptide by a variety of recombinant techniques. The molecule is useful for generating new cloning and transformed and transfected expression vectors, prokaryotic and eukaryotic host cells, and new and useful methods for cultured growth of such host cells capable of expression of the polypeptide and related products.

Moreover, the isolated mammalian nucleic acid molecules encoding a mammalian prostate-specific membrane antigen and the alternatively spliced PSM' are useful for the development of probes to study the tumorigenesis of

prostate cancer.

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This invention also provides an isolated nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule encoding the prostate-specific membrane antigen or the alternatively spliced prostate specific membrane antigen.

This nucleic acid molecule produced can either be DNA or RNA. As used herein, the phrase "specifically hybridizing" means the ability of a nucleic acid molecule to recognize a nucleic acid sequence complementary to its own and to form double-helical segments through hydrogen bonding between complementary base pairs.

This nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule encoding the prostate-specific membrane antigen can be used as a probe. Nucleic acid probe technology is well known to those skilled in the art who will readily appreciate that such probes may vary greatly in length and may be labeled with a detectable label, such as a radioisotope or fluorescent dye, to facilitate detection of the probe. DNA probe molecules may be produced by insertion of a DNA molecule which encodes PSM antigen into suitable vectors, such as plasmids or bacteriophages, followed by transforming into suitable bacterial host cells, replication in the transformed bacterial host cells and harvesting of the DNA probes, using methods well known in the art. Alternatively, probes may be generated chemically from DNA synthesizers.

RNA probes may be generated by inserting the PSM antigen molecule downstream of a bacteriophage promoter

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such as T3, T7 or SP6. Large amounts of RNA probe may be produced by incubating the labeled nucleotides with the linearized PSM antigen fragment where it contains an upstream promoter in the presence of the appropriate RNA polymerase.

This invention also provides a nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of a nucleic acid molecule which is complementary to the mammalian nucleic acid molecule encoding a mammalian prostate-specific membrane antigen. This molecule may either be a DNA or RNA molecule.

The current invention further provides a method of 15 detecting the expression of a mammalian PSM or PSM' antigen expression in a cell which comprises obtaining total mRNA from the cell, contacting the mRNA so obtained with a labelled nucleic acid molecule of at specifically nucleotides capable of 15 20 hybridizing with a sequence of the nucleic acid molecule encoding a mammalian PSM or PSM' antigen under hybridizing conditions, determining the presence of mRNA hybridized to the molecule and thereby detecting the expression of the mammalian prostate-specific 25 membrane antigen in the cell. The nucleic acid molecules synthesized above may be used to detect expression of a PSM or PSM' antigen by detecting the presence of mRNA coding for the PSM antigen. mRNA from the cell may be isolated by many procedures 30 well known to a person of ordinary skill in the art. The hybridizing conditions of the labelled nucleic acid molecules may be determined by routine experimentation well known in the art. The presence of mRNA hybridized to the probe may be determined by gel electrophoresis 35 or other methods known in the art. By measuring the amount of the hybrid made, the expression of the PSM antigen by the cell can be determined. The labeling For an example, one or more may be radioactive. radioactive nucleotides can be incorporated in the nucleic acid when it is made.

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In one embodiment of this invention, nucleic acids are extracted by precipitation from lysed cells and the mRNA is isolated from the extract using an oligo-dT column which binds the poly-A tails of the mRNA then exposed is mRNA molecules (13). The radioactively labelled probe on a nitrocellulose membrane, and the probe hybridizes to and thereby Binding may be labels complementary mRNA sequences. autoradiography luminescence by detected However, other methods for scintillation counting. performing these steps are well known to those skilled in the art, and the discussion above is merely an example.

This invention further provides another method to 20 detect expression of a PSM or PSM' antigen in tissue sections which comprises contacting the tissue sections with a labelled nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of nucleic acid molecules encoding a mammalian 25 PSM antigen under hybridizing conditions, determining the presence of mRNA hybridized to the molecule and thereby detecting the expression of the mammalian PSM or PSM' antigen in tissue sections. The probes are also useful for in-situ hybridization or in order to 30 locate tissues which express this gene, or for other hybridization assays for the presence of this gene or its mRNA in various biological tissues. hybridization using a labelled nucleic acid molecule is well known in the art. Essentially, tissue sections 35 are incubated with the labelled nucleic acid molecule to allow the hybridization to occur. The molecule will

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carry a marker for the detection because it is "labelled", the amount of the hybrid will be determined based on the detection of the amount of the marker and so will the expression of PSM antigen.

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This invention further provides isolated PSM or PSM' antigen nucleic acid molecule operatively linked to a promoter of RNA transcription. The isolated PSM or PSM' antigen sequence can be linked to vector systems. Various vectors including plasmid vectors, cosmid vectors, bacteriophage vectors and other viruses are well known to ordinary skilled practitioners. This invention further provides a vector which comprises the isolated nucleic acid molecule encoding for the PSM or PSM' antigen.

As an example to obtain these vectors, insert and vector DNA can both be exposed to a restriction enzyme to create complementary ends on both molecules which base pair with each other and are then ligated together with DNA ligase. Alternatively, linkers can be ligated to the insert DNA which correspond to a restriction site in the vector DNA, which is then digested with the restriction enzyme which cuts at that site. Other means are also available and known to an ordinary skilled practitioner.

In an embodiment, the PSM sequence is cloned in the Not I/Sal I site of pSPORT/vector (Gibco® - BRL). plasmid, p55A-PSM, was deposited on August 14, 1992 30 with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A. under the provisions of the Budapest Treaty for Deposit the of International Recognition Microorganism for the Purposes of Patent Procedure. 35 Plasmid, p55A-PSM, was accorded ATCC Accession Number 75294.

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This invention further provides a host vector system for the production of a polypeptide having the biological activity of the prostate-specific membrane antigen. These vectors may be transformed into a suitable host cell to form a host cell vector system for the production of a polypeptide having the biological activity of PSM antigen.

Regulatory elements required for expression include promoter sequences to bind RNA polymerase 10 sequences for ribosome initiation transcription For example, a bacterial expression vector includes a promoter such as the lac promoter and for transcription initiation the Shine-Dalgarno sequence and the start codon AUG (14). Similarly, a eukaryotic 15 expression vector includes a heterologous or homologous II, а downstream . RNA polymerase for polyadenylation signal, the start codon AUG, and a termination codon for detachment of the ribosome. Such vectors may be obtained commercially or assembled from 20 the sequences described by methods well known in the art, for example the methods described above for constructing vectors in general. Expression vectors are useful to produce cells that express the PSM antigen. 25

This invention further provides an isolated DNA or cDNA molecule described hereinabove wherein the host cell is selected from the group consisting of bacterial cells (such as <u>E.coli</u>), yeast cells, fungal cells, insect cells and animal cells. Suitable animal cells include, but are not limited to Vero cells, HeLa cells, Cos cells, CV1 cells and various primary mammalian cells.

This invention further provides a method of producing a polypeptide having the biological activity of the prostate-specific membrane antigen which comprising

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growing host cells of a vector system containing the PSM antigen sequence under suitable conditions permitting production of the polypeptide and recovering the polypeptide so produced.

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This invention provides a mammalian cell comprising a DNA molecule encoding a mammalian PSM or PSM' antigen, such as a mammalian cell comprising a plasmid adapted for expression in a mammalian cell, which comprises a DNA molecule encoding a mammalian PSM antigen and the regulatory elements necessary for expression of the DNA in the mammalian cell so located relative to the DNA encoding the mammalian PSM or PSM' antigen as to permit expression thereof.

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Numerous mammalian cells may be used as hosts, including, but not limited to, the mouse fibroblast cell NIH3T3, CHO cells, HeLa cells, Ltk cells, Cos cells, etc. Expression plasmids such as that described supra may be used to transfect mammalian cells by methods well known in the art such as calcium phosphate precipitation, electroporation or DNA encoding the mammalian PSM antigen may be otherwise introduced into mammalian cells, e.g., by microinjection, to obtain mammalian cells which comprise DNA, e.g., cDNA or a plasmid, encoding a mammalian PSM antigen.

This invention provides a method for determining whether a ligand can bind to a mammalian prostate-specific membrane antigen which comprises contacting a mammalian cell comprising an isolated DNA molecule encoding a mammalian prostate-specific membrane antigen with the ligand under conditions permitting binding of ligands to the mammalian prostate-specific membrane antigen, and thereby determining whether the ligand binds to a mammalian prostate-specific membrane antigen.

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This invention further provides ligands bound to the mammalian PSM or PSM' antigen.

This invention also provides a therapeutic agent comprising a ligand identified by the above-described method and a cytotoxic agent conjugated thereto. The cytotoxic agent may either be a radioisotope or a toxin. Examples of radioisotopes or toxins are well known to one of ordinary skill in the art.

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This invention also provides a method of imaging prostate cancer in human patients which comprises administering to the patients at least one ligand identified by the above-described method, capable of binding to the cell surface of the prostate cancer cell and labelled with an imaging agent under conditions permitting formation of a complex between the ligand and the cell surface PSM or PSM' antigen. invention further provides a composition comprising an effective imaging agent of the PSM OR PSM' antigen ligand and a pharmaceutically acceptable carrier. Pharmaceutically acceptable carriers are well known to one of ordinary skill in the art. For an example, such acceptable carrier pharmaceutically physiological saline.

Also provided by this invention is a purified mammalian As used herein, the term PSM and PSM' antigen. "purified prostate-specific membrane antigen" shall mean isolated naturally-occurring prostate-specific membrane antigen or protein (purified from nature or manufactured such that the primary, secondary and posttranslational and conformation, tertiary modifications are identical to naturally-occurring non-naturally occurring as well as polypeptides having a primary structural conformation (i.e. continuous sequence of amino acid residues).

Such polypeptides include derivatives and analogs.

This invention provides an isolated nucleic acid molecule encoding a prostate-specific membrane antigen promoter. In one embodiment the PSM promoter has at least the sequence as in Figures 58A-58C.

This invention provides an isolated nucleic acid molecule encoding an alternatively spliced prostate-specific membrane antigen promoter.

This invention further provides a polypeptide encoded by the isolated mammalian nucleic acid sequence of PSM and PSM' antigen.

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is believed that there may be natural ligand interacting with the PSM or PSM' antigen. This invention provides a method to identify such natural ligand or other ligand which can bind to the PSM or antigen. A method to identify the ligand comprises a) coupling the purified mammalian PSM or PSM' antigen to a solid matrix, b) incubating the coupled purified mammalian PSM or PSM' protein with the potential ligands under the conditions permitting binding of ligands and the purified PSM or PSM' antigen; c) washing the ligand and coupled purified mammalian PSM or PSM' antigen complex formed in b) to eliminate the nonspecific binding and impurities and finally d) eluting the ligand from the bound purified mammalian PSM or PSM' antigen. The techniques of coupling proteins to a solid matrix are well known in the art. Potential ligands may either be deduced from the structure of mammalian PSM or PSM' by other empirical experiments known by ordinary The conditions for binding may also practitioners. easily be determined and protocols for carrying such experimentation have long been well documented (15).

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The ligand-PSM antigen complex will be washed. Finally, the bound ligand will be eluted and characterized. Standard ligands characterization techniques are well known in the art.

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The above method may also be used to purify ligands from any biological source. For purification of natural ligands in the cell, cell lysates, serum or other biological samples will be used to incubate with the mammalian PSM or PSM' antigen bound on a matrix. Specific natural ligand will then be identified and purified as above described.

with the protein sequence information, antigenic areas may be identified and antibodies directed against these areas may be generated and targeted to the prostate cancer for imaging the cancer or therapies.

This invention provides an antibody directed against the amino acid sequence of a mammalian PSM or PSM' antigen.

This invention provides a method to select specific regions on the PSM or PSM' antigen to generate The protein sequence may be determined antibodies. from the PSM or PSM' DNA sequence. Amino acid sequences may be analyzed by methods well known to those skilled in the art to determine whether they produce hydrophobic or hydrophilic regions in the In the case of cell proteins which they build. membrane proteins, hydrophobic regions are well known to form the part of the protein that is inserted into lipid bilayer of the cell membrane, hydrophilic regions are located on the cell surface, in Usually, the hydrophilic an aqueous environment. regions will be more immunogenic than the hydrophobic Therefore the hydrophilic amino acid regions.

sequences may be selected and used to generate antibodies specific to mammalian PSM antigen. For an example, hydrophilic sequences of the human PSM antigen shown in hydrophilicity plot of Figures 16:1-11 may be easily selected. The selected peptides may be prepared using commercially available machines. As an alternative, DNA, such as a cDNA or a fragment thereof, may be cloned and expressed and the resulting polypeptide recovered and used as an immunogen.

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Polyclonal antibodies against these peptides may be produced by immunizing animals using the selected peptides. Monoclonal antibodies are prepared using hybridoma technology by fusing antibody producing B cells from immunized animals with myeloma cells and selecting the resulting hybridoma cell line producing the desired antibody. Alternatively, monoclonal antibodies may be produced by in vitro techniques known to a person of ordinary skill in the art. These antibodies are useful to detect the expression of mammalian PSM antigen in living animals, in humans, or in biological tissues or fluids isolated from animals or humans.

- In one embodiment, peptides Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. ), Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID No. ) and Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID No. ) of human PSM antigen are selected.
- This invention further provides polyclonal and monoclonal antibody(ies) against peptides Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. ), Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID No. ) and Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID No. ).

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This invention provides a therapeutic agent comprising antibodies or ligand(s) directed against PSM antigen

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and a cytotoxic agent conjugated thereto or antibodies linked enzymes which activate prodrug to kill the tumor. The cytotoxic agent may either be a radioisotope or toxin.

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This invention provides a method of imaging prostate cancer in human patients which comprises administering to the patient the monoclonal antibody directed against the peptide of the mammalian PSM or PSM' antigen capable of binding to the cell surface of the prostate cancer cell and labeled with an imaging agent under conditions permitting formation of a complex between the monoclonal antibody and the cell surface prostate-specific membrane antigen. The imaging agent is a radioisotope such as Indium<sup>111</sup>.

This invention further provides a prostate cancer specific imaging agent comprising the antibody directed against PSM or PSM' antigen and a radioisotope conjugated thereto.

This invention also provides a composition comprising an effective imaging amount of the antibody directed against the PSM or PSM' antigen and a pharmaceutically acceptable carrier. The methods to determine effective imaging amounts are well known to a skilled practitioner. One method is by titration using different amounts of the antibody.

This invention further provides an immunoassay for measuring the amount of the prostate-specific membrane antigen in a biological sample comprising steps of a) contacting the biological sample with at least one antibody directed against the PSM or PSM' antigen to form a complex with said antibody and the prostate-specific membrane antigen, and b) measuring the amount of the prostate-specific membrane antigen in said

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biological sample by measuring the amount of said complex. One example of the biological sample is a serum sample.

This invention provides a method to purify mammalian prostate-specific membrane antigen comprising steps of a) coupling the antibody directed against the PSM or PSM' antigen to a solid matrix; b) incubating the coupled antibody of a) with lysate containing prostate-specific membrane antigen under the condition which the antibody and prostate membrane specific can bind; c) washing the solid matrix to eliminate impurities and d) eluting the prostate-specific membrane antigen from the coupled antibody.

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This invention also provides a transgenic nonhuman mammal which comprises the isolated nucleic acid molecule encoding a mammalian PSM or PSM' antigen. This invention further provides a transgenic nonhuman mammal whose genome comprises antisense DNA complementary to DNA encoding a mammalian prostate-specific membrane antigen so placed as to be transcribed into antisense mRNA complementary to mRNA encoding the prostate-specific membrane antigen and which hybridizes to mRNA encoding the prostate specific antigen thereby reducing its translation.

Animal model systems which elucidate the physiological and behavioral roles of mammalian PSM or PSM' antigen are produced by creating transgenic animals in which the expression of the PSM or PSM' antigen is either increased or decreased, or the amino acid sequence of the expressed PSM antigen is altered, by a variety of techniques. Examples of these techniques include, but are not limited to: 1) Insertion of normal or mutant versions of DNA encoding a mammalian PSM or PSM' antigen, by microinjection, electroporation, retroviral

transfection or other means well known to those skilled in the art, into appropriate fertilized embryos in order to produce a transgenic animal (16) or Homologous recombination (17) of mutant or normal, human or animal versions of these genes with the native locus in transgenic animals to alter regulation of expression or the structure of these PSM or PSM' antigen sequences. The technique of homologous recombination is well known in the art. It replaces the native gene with the inserted gene and so is useful for producing an animal that cannot express native PSM antigen but does express, for example, an inserted mutant PSM antigen, which has replaced the native PSM antigen in the animal's genome by recombination, resulting in undere xpression of the transporter. Microinjection adds genes to the genome, but does not remove them, and so is useful for producing an animal which expresses its own and added PSM antigens, resulting in over expression of the PSM antigens.

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One means available for producing a transgenic animal, with a mouse as an example, is as follows: mice are mated, and the resulting fertilized eggs are The eggs are stored dissected out of their oviducts. in an appropriate medium such as Me medium (16). DNA or cDNA encoding a mammalian PSM antigen is purified from a vector by methods well known in the art. Inducible promoters may be fused with the coding region of the DNA to provide an experimental means to regulate Alternatively or in expression of the trans-gene. addition, tissue specific regulatory elements may be fused with the coding region to permit tissue-specific expression of the trans-gene. The DNA, in appropriately buffered solution, put into is microinjection needle (which may be made from capillary tubing using a pipet puller) and the egg to be injected is put in a depression slide. The needle is inserted

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into the pronucleus of the egg, and the DNA solution is injected. The injected egg is then transferred into the oviduct of a pseudopregnant mouse (a mouse stimulated by the appropriate hormones to maintain pregnancy but which is not actually pregnant), where it proceeds to the uterus, implants, and develops to term. As noted above, microinjection is not the only method for inserting DNA into the egg cell, and is used here only for exemplary purposes.

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Another use of the PSM antigen sequence is to isolate homologous gene or genes in different mammals. The gene or genes can be isolated by low stringency screening of either cDNA or genomic libraries of different mammals using probes from PSM sequence. The positive clones identified will be further analyzed by DNA sequencing techniques which are well known to an ordinary person skilled in the art. For example, the detection of members of the protein serine kinase family by homology probing.

This invention provides a method of suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells comprising introducing a DNA molecule encoding a prostate specific membrane antigen operatively linked to a 5' regulatory element into a tumor cell of a subject, in a way that expression of the prostate specific membrane antigen is under the control of the regulatory element, thereby suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells. The subject may be a mammal or more specifically a human.

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In one embodiment, the DNA molecule encoding prostate specific membrane antigen operatively linked to a 5'

regulatory element forms part of a transfer vector which is inserted into a cell or organism. In addition the vector is capable or replication and expression of prostate specific membrane antigen. The DNA molecule encoding prostate specific membrane antigen can be integrated into a genome of a eukaryotic or prokaryotic cell or in a host cell containing and/or expressing a prostate specific membrane antigen.

Further, the DNA molecule encoding prostate specific membrane antigen may be introduced by a bacterial, viral, fungal, animal, or liposomal delivery vehicle. Other means are also available and known to an ordinary skilled practitioner.

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Further, the DNA molecule encoding a prostate specific membrane antigen operatively linked to a promoter or enhancer. A number of viral vectors have been described including those made from various promoters and other regulatory elements derived from virus sources. Promoters consist of short arrays of nucleic acid sequences that interact specifically with cellular proteins involved in transcription. The combination of different recognition sequences and the cellular concentration of the cognate transcription factors determines the efficiency with which a gene is transcribed in a particular cell type.

Examples of suitable promoters include a viral promoter. Viral promoters include: adenovirus promoter, an simian virus 40 (SV40) promoter, a cytomegalovirus (CMV) promoter, a mouse mammary tumor virus (MMTV) promoter, a Malony murine leukemia virus promoter, a murine sarcoma virus promoter, and a Rous sarcoma virus promoter.

Further, another suitable promoter is a heat shock

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promoter. Additionally, a suitable promoter is a bacteriophage promoter. Examples of suitable bacteriophage promoters include but not limited to, a T7 promoter, a T3 promoter, an SP6 promoter, a lambda promoter, a baculovirus promoter.

Also suitable as a promoter is an animal cell promoter such as an interferon promoter, a metallothionein promoter, an immunoglobulin promoter. A fungal promoter is also a suitable promoter. Examples of fungal promoters include but are not limited to, an ADC1 promoter, an ARG promoter, an ADH promoter, a CYC1 promoter, a CUP promoter, an ENO1 promoter, a GAL promoter, a PHO promoter, a PGK promoter, a GAPDH promoter, a mating type factor promoter. Further, plant cell promoters and insect cell promoters are also suitable for the methods described herein.

This invention provides a method of suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells, comprising introducing a DNA molecule encoding

prostate specific membrane antigen operatively linked to a 5' regulatory element coupled with a therapeutic DNA into a tumor cell of a subject, thereby suppressing or modulating metastatic ability of prostate tumor cells, prostate tumor growth or elimination of prostate tumor cells. The subject may be a mammal or more specifically a human.

Further, the therapeutic DNA which is coupled to the DNA molecule encoding a prostate specific membrane antigen operatively linked to a 5' regulatory element into a tumor cell may code for a cytokine, viral antigen, or a pro-drug activating enzyme. Other means are also available and known to an ordinary skilled

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practitioner.

In addition, this invention provides a prostate tumor cell, comprising a DNA molecule isolated from mammalian nucleic acid encoding a mammalian prostate-specific membrane antigen under the control of a prostate specific membrane antigen operatively linked to a 5' regulatory element.

As used herein, DNA molecules include complementary DNA (cDNA), synthetic DNA, and genomic DNA.

This invention provides a therapeutic vaccine for preventing human prostate tumor growth or stimulation of prostate tumor cells in a subject, comprising administering an effective amount to the prostate cell, and a pharmaceutical acceptable carrier, thereby preventing the tumor growth or stimulation of tumor cells in the subject. Other means are also available and known to an ordinary skilled practitioner.

method of detecting provides a This invention hematogenous micrometastic tumor cells of a subject, comprising (A) performing nested polymerase chain reaction (PCR) on blood, bone marrow or lymph node samples of the subject using the prostate specific membrane antigen primers or alternatively spliced prostate specific antigen primers, and (B) verifying sequencing Southern and micrometastases by DNA analysis, thereby detecting hematogenous micrometastic tumor cells of the subject. The subject may be a mammal or more specifically a human.

The micrometastatic tumor cell may be a prostatic cancer and the DNA primers may be derived from prostate specific antigen. Further, the subject may be administered with simultaneously an effective amount of

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hormones, so as to increase expression of prostate specific membrane antigen. Further, growth factors or cytokine may be administered in separately or in conjunction with hormones. Cytokines include, but are not limited to: transforming growth factor beta, epidermal growth factor (EGF) family, fibroblast growth factors, hepatocyte growth factor, insulin-like growth factors, B-nerve growth factor, platelet-derived growth factor, vascular endothelial growth factor, interleukin 1, IL-1 receptor antagonist, interleukin 2, interleukin 3, interleukin 4, interleukin 5, interleukin 6, IL-6 interleukin 7, interleukin soluble receptor, interleukin 11. interleukin 9. interleukin 10, interleukin 12, interleukin 13, angiogenin, chemokines, colony stimulating factors, granulocyte-macrophage colony stimulating factors, erythropoietin, interferon, inhibitory leukemia gamma, interferon secretory leukocyte Μ, pleiotrophin, oncostatin protease inhibitor, stem cell factor, tumor necrosis factors, adhesion molecule, and soluble tumor necrosis factor (TNF) receptors.

This invention provides a method of abrogating the mitogenic response due to transferrin, comprising introducing a DNA molecule encoding prostate specific membrane antigen operatively linked to a 5' regulatory element into a tumor cell, the expression of which gene is directly associated with a defined pathological effect within a multicellular organism, thereby abrogating mitogen response due to transferrin. The tumor cell may be a prostate cell.

This invention provides a method of determining prostate cancer progression in a subject which comprises: a) obtaining a suitable prostate tissue sample; b) extracting RNA from the prostate tissue sample; c) performing a RNAse protection assay on the

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RNA thereby forming a duplex RNA-RNA hybrid; d) detecting PSM and PSM' amounts in the tissue sample; e) calculating a PSM/PSM' tumor index, thereby determining prostate cancer progression in the subject. In-situ hyribridization may be performed in conjunction with the above detection method.

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This invention provides a method of detecting prostate cancer in a subject which comprises: (a) obtaining from a subject a prostate tissue sample; (b) treating the tissue sample so as to separately recover nucleic acid molecules present in the prostate tissue sample; (c) contacting the resulting nucleic acid molecules single-stranded multiple pairs of oligonucleotide primers, each such pair being capable of specifically hybridizing to the tissue sample, under hybridizing conditions; (d) amplifying any nucleic acid molecules to which a pair of primers hybridizes so as to obtain a double-stranded amplification product; (e) treating any such double-stranded amplification product so as to obtain single-stranded nucleic acid (f) contacting any resulting molecules therefrom; single-stranded nucleic acid molecules with multiple single-stranded labeled oligonucleotide probes, each such probe containing the same label and being capable of specifically hybridizing with such tissue sample, (g) contacting any under hybridizing conditions; resulting hybrids with an antibody to which a marker is attached and which is capable of specifically forming a complex with the labeled-probe, when the probe is 30 present in such a complex, under complexing conditions; and (h) detecting the presence of any resulting complexes, the presence thereof being indicative of prostate cancer in a subject.

This invention provides a method of enhancing antibody based targeting of PSM or PSM' in prostate tissue for more hour intervals by a subsequent injection or other administration.

As used herein administration means a method of administering to a subject. Such methods are well known to those skilled in the art and include, but are not limited to, administration topically, parenterally, orally, intravenously, intramuscularly, subcutaneously or by aerosol. Administration of PSM may be effected continuously or intermittently.

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The pharmaceutical formulations or compositions of this invention may be in the dosage form of solid, semisolid, or liquid such as, e.g., suspensions, aerosols Preferably the compositions the like. administered in unit dosage forms suitable for single administration of precise dosage amounts. The compositions may also include, depending on the formulation desired, pharmaceutically-acceptable, nontoxic carriers or diluents, which are defined as vehicles commonly used to formulate pharmaceutical compositions for animal or human administration. diluent is selected so as not to affect the biological activity of the combination. Examples of such diluents are distilled water, physiological saline, Ringer's solution, dextrose solution, and Hank's solution. addition, the pharmaceutical composition or formulation may also include other carriers, adjuvants; nontoxic, nontherapeutic, nonimmunogenic stabilizers Effective amounts of such diluent or and the like. carrier are those amounts which are effective to obtain a pharmaceutically acceptable formulation in terms of solubility of components, or biological activity, etc

This invention will be better understood from the Experimental Details which follow. However, one skilled in the art will readily appreciate that the

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specific methods and results discussed are merely illustrative of the invention as described more fully in the claims which follow thereafter.

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### EXPERIMENTAL DETAILS

### EXAMPLE 1:

Materials and Methods: The approach for cloning the 5 involved purification of antigen the immunoprecipitation, and microsequencing of several internal peptides for use in synthesizing degenerate oligonucleotide primers for subsequent use in the polymerase chain reaction (19, 20). A partial cDNA was 10 amplified as a PCR product and this was used as a homologous probe to clone the full-length cDNA molecule from a LNCaP (Lymph Node Carcinoma of Prostate) cell line cDNA plasmid library (8).

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Western Analysis of the PSM Antigen: Membrane proteins were isolated from cells by hypotonic lysis followed by centrifugation over a sucrose density gradient (21).  $10-20\mu g$  of LNCaP, DU-145, and PC-3 membrane proteins were electrophoresed through a 10% SDS-PAGE resolving 20 gel with a 4% stacking gel at 9-10 milliamps for 16-18 Proteins were electroblotted onto PVDF membranes (Millipore® Corp.) in transfer buffer (48mM Tris base, 39mM Glycine, 20% Methanol) at 25 volts overnight at 4°C. Membranes were blocked in TSB (0.15M 25 NaCl, 0.01M Tris base, 5% BSA) for 30 minutes at room temperature followed by incubation with  $10-15\mu g/ml$  of CYT-356 monoclonal antibody (Cytogen Corp.) for 2 Membranes were then incubated with  $10-15\mu g/ml$ hours. immunoglobulin (Accurate anti-mouse rabbit 30 Scientific) for 1 hour at room temperature followed by incubation with 125I-Protein A (Amersham®), at 1x106 cpm/ml at room temperature. Membranes were then washed and autoradiographed for 12-24 hours at -70°C (Figure 1). 35

Immunohistochemical Analysis of PSM Antigen Expression: method avidin-biotin of immunohistochemical detection was employed to analyze both human tissue sections and cell lines for PSM Antigen expression Cryostat-cut prostate tissue sections (4-6µm 5 thick) were fixed in methanol/acetone for 10 minutes. Cell cytospins were made on glass slides using 50,000 cells/100µl/slide. Samples were treated with 1% hydrogen peroxide in PBS for 10-15 minutes in order to remove any endogenous peroxidase activity. 10 sections were washed several times in PBS, and then incubated with the appropriate suppressor serum for 20 minutes. The suppressor serum was drained off and the sections or cells were then incubated with the diluted CYT-356 monoclonal antibody for 1 hour. Samples were 15 then washed with PBS and sequentially incubated with secondary antibodies (horse or goat immunoglobulins, 1:200 dilution for 30 minutes), and with avidin-biotin complexes (1:25 dilution for 30 minutes). DAB was used as a chromogen, followed by hematoxylin counterstaining 20 and mounting. Frozen sections of prostate samples and duplicate cell cytospins were used as controls for each a positive control, experiment. As the anticytokeratin monoclonal antibody CAM 5.2 was used following the same procedure described above. 25 sections are considered by us to express the PSM antigen if at least 5% of the cells demonstrate immunoreactivity. The scoring system is as follows: 1 = <5%; 2 = 5-19%; 3 = 20-75%; and 4 = >75% positive cells. Homogeneity versus heterogeneity was accounted 30 for by evaluating positive and negative cells in 3-5 high power light microscopic fields (400x), recording the percentage of positive cells among 100-500 cells. The intensity of immunostaining is graded on a 1+ to 4+ 35 scale, where 1+ represents mild, 2-3+ represents moderate, and 4+ represents intense immunostaining as compared to positive controls.

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Immunoprecipitation of the PSM Antigen: 80%-confluent LNCaP cells in 100mm petri dishes were starved in RPMI media without methionine for 2 hours, after which 35S-Methionine was added at  $100\mu\text{Ci/ml}$  and the cells were grown for another 16-18 hours. Cells were then washed and lysed by the addition of 1ml of lysis buffer (1% Triton X-100, 50mM Hepes pH 7.5, 10% glycerol, 150mM MgCl, 1mM PMSF, and 1mM EGTA) with incubation for 20 minutes at 4°C. Lysates were pre-cleared by mixing with Pansorbin® cells (Calbiochem®) for 90 minutes at Cell lysates were then mixed with Protein A Sepharose® CL-4B beads (Pharmacia®) previously bound with CYT-356 antibody (Cytogen Corp.) and RAM antibody (Accurate Scientific) for 3-4 hours at 4°C. antibody was used per 3mg of beads per petri dish. Beads were then washed with HNTG buffer (20mM Hepes pH 7.5, 150mM NaCl, 0.1% Triton X-100, 10% glycerol, and 2mM Sodium Orthovanadate), resuspended in sample loading buffer containing &-mercaptoethanol, denatured at 95°C for 5-10 minutes and run on a 10% SDS-PAGE gel with a 4° stacking gel at 10 milliamps overnight. Gels were stained with Coomassie Blue, destained with acetic acid/methanol, and dried down in a vacuum dryer at 60°C. Gels were then autoradiographed for 16-24 hours at -70°C (Figures 2A-2D).

# Immunoprecipitation and Peptide Sequencing:

The procedure described above for immunoprecipitation was repeated with 8 confluent petri dishes containing The LNCaP 6x10<sup>7</sup> approximately immunoprecipitation product was pooled and loaded into two lanes of a 10% SDS-PAGE gel and electrophoresed at Proteins hours. milliamps for 16 electroblotted onto Nitrocellulose BA-85 membranes (Schleicher and Schuell®) for 2 hours at 75 volts at 4°C in transfer buffer. Membranes were stained with Ponceau Red to visualize the proteins and the 100kD

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protein band was excised, solubilized, and digested proteolytically with trypsin. HPLC was then performed on the digested sample on an Applied Biosystems Model 171C and clear dominant peptide peaks were selected and sequenced by modified Edman degradation on a modified Biosystems liquid Applied Model Protein/Peptide Microsequencer (23). Sequencing data on all of the peptides is included within this The amino-terminus of the PSM antigen was sequenced by a similar method which involved purifying the antigen by immunoprecipitation and transfer via electro-blotting to a PVDF membrane (Millipore®). Protein was analyzed on an Applied Biosystems Model 477A Protein/Peptide Sequencer and the amino terminus was found to be blocked, and therefore no sequence data could be obtained by this technique.

#### PSM Antigen Peptide Sequences:

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      2T17 #5
                 SLYES(W) TK (SEQ ID No.
                 (S) YPDGXNLPGG (g) VQR (SEQ ID No. )
      2T22 #9
      2T26 #3
                 FYDPMFK (SEQ ID No.
      2T27 #4
                 IYNVIGTL(K) (SEQ ID No.
      2T34 #6
                 FLYXXTQIPHLAGTEQNFQLAK (SEQ ID No. )
      2T35 #2
                 G/PVILYSDPADYFAPD/GVK (SEQ ID No. )
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      2T38 #1
                AFIDPLGLPDRPFYR (SEQ ID No.
                 YAGESFPGIYDALFDIESK (SEQ ID No.
      2T46 #8
                 TILFAS (W) DAEEFGXX (q) STE (e) A (E) ... (SEQ ID No.
      2T47 #7
       )
```

Notes: X means that no residue could be identified at this position. Capital denotes identification but with a lower degree of confidence. (lower case) means residue present but at very low levels. ... indicates sequence continues but has dropped below detection limit.

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All of these peptide sequences were verified to be unique after a complete homology search of the translated Genbank computer database.

Degenerate PCR: Sense and anti-sense 5'unphosphorylated degenerate oligonucleotide primers 17
to 20 nucleotides in length corresponding to portions
of the above peptides were synthesized on an Applied
Biosystems Model 394A DNA Synthesizer. These primers
have degeneracies from 32 to 144. The primers used are
shown below. The underlined amino acids in the
peptides represent the residues used in primer design.

### Peptide 3: FYDPMFK (SEQ ID No. )

PSM Primer "A" TT(C or T) - TA(C or T) - GA(C or T) - CCX - ATG - TT (SEQ ID No. )

PSM Primer "B" AAC - ATX - GG(A or G) - TC(A or G) 
TA(A or G) - AA (SEQ ID No. )

Primer A is sense primer and B is anti-sense. Degeneracy is 32-fold.

25 Peptide 4: IYNVIGTL(K) (SEQ ID No. 6)

PSM Primer "C" AT(T or C or A) - TA(T or C) - AA(T or C) - GTX - AT(T or C or A) - GG (SEQ ID No.)

PSM Primer "D" CC(A or T or G) - ATX - AC(G or A) - TT(A or G) - TA(A or G or T) - AT (SEQ ID No.)

Primer C is sense primer and D is anti-sense. Degeneracy is 144-fold.

Peptide 2: G/PVILYSDPADYFAPD/GVK (SEQ ID No. )

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PSM Primer "E" CCX - GCX - GA(T or C) - TA(T or C) - TT(T or C) - GC (SEQ ID No. )
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PSM Primer "F" GC(G or A) - AA(A or G) - TA(A or G) 
TXC - GCX - GG (SEQ ID No. )

Primer E is sense primer and F is antisense primer. Degeneracy is 128-fold.

10 Peptide 6: FLYXXTQIPHLAGTEONFOLAK (SEQ ID No. )

PSM Primer "I" ACX - GA(A or G) - CA(A or G) - AA(T or C) - TT(T or C) - CA(A or G) - CT (SEQ ID No. )

PSM Primer "J" AG - (T or C)TG - (A or G)AA - (A or G)TT - (T or C)TG - (T or C)TC - XGT (SEQ ID No. )

PSM Primer "K" GA(A or G) - CA(A or G) - AA(T or C) - TT(T or C) CA(A or G) - CT (SEQ ID No. )

PSM Primer "L" AG - (T or C)TG - (A or G)AA - (A or G)TT - (T or C)TG - (T or C)TC (SEQ ID No. 22)

Primers I and K are sense primers and J and L are antisense. I and J have degeneracies of 128-fold and K and L have 32-fold degeneracy.

Peptide 7: TILFAS (W) DAEEFGXX (q) STE (e) A (E) ... (SEQ ID No. )

PSM Primer "M" TGG - GA(T or C) - GCX - GA(A or G) - GA(A or G) - TT(C or T) - GG (SEQ ID No. )

PSM Primer "N" CC - (G or A)AA - (T or C)TC - (T or 35 C)TC - XGC - (A or G)TC - CCA (SEQ ID No.)

PSM Primer "O" TGG - GA(T or C) - GCX - GA(A or G) -

GA(A or G) - TT (SEQ ID No. )

PSM Primer "P" AA - (T or C)TC - (T or C)TC - XGC - (A or G)TC - CCA (SEQ ID No. )

5 Primers M and O are sense primers and N and P are antisense. M and N have degeneracy of 64-fold and O and P are 32-fold degenerate.

Degenerate PCR was performed using a Perkin-Elmer Model 10 480 DNA thermal cycler. cDNA template for the PCR was prepared from LNCaP mRNA which had been isolated by chromatography dT oligo methods of standard (Collaborative Research). The cDNA synthesis was carried out as follows:

> LNCaP poly A+ RNA  $(2\mu g)$  $4.5\mu$ l

Oligo dT primers (0.5µg) 1.0µl

<u>dH,O</u> <u>4:5µl</u>

10µl 20

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Incubate at 68°C x 10 minutes. Quick chill on ice x 5 minutes.

#### Add: 25

5 x RT Buffer  $4\mu l$ 

0.1M DTT  $2\mu l$ 

10mM dNTPs  $1\mu l$ 

RNasin (Promega)  $0.5 \mu l$ 30

> <u>dH</u>,0  $1.5\mu$ l

 $19\mu$ l

Incubate for 2 minutes at 37°C.

Add 1µl Superscript® Reverse Transcriptase (Gibco®-BRL) 35 Incubate for 1 hour at 37°C.

Add  $30\mu l$  dH<sub>2</sub>O. Use  $2\mu l$  per PCR reaction.

Degenerate PCR reactions were optimized by varying the annealing temperatures, Mg++ concentrations, primer concentrations, buffer composition, extension times and number of cycles. The optimal thermal cycler profile was: Denaturation at 94°C x 30 seconds, Annealing at 45-55°C for 1 minute (depending on the mean T<sub>m</sub> of the primers used), and Extension at 72°C for 2 minutes.

10 x PCR Buffer\* 5µ1 2.5mM dNTP Mix 5µl Primer Mix (containing 0.5-1.0µg each of 5µ1 and anti-sense primers) sense 15 100mM ß-mercaptoethanol 5ul  $2\mu l$ LNCaP cDNA template 25mM MgCl, (2.5mM final) 5µ1 21µ1 dH,O diluted Taq Polymerase  $(0.5U/\mu l)$ 20  $2\mu l$  $50\mu$ l total volume

Tubes were overlaid with  $60\mu$ l of light mineral oil and amplified for 30 cycles. PCR products were analyzed by electrophoresing  $5\mu$ l of each sample on a 2-3% agarose gel followed by staining with Ethidium bromide and photography.

## \*10x PCR Buffer 166mM NH<sub>4</sub>SO<sub>4</sub> 670mM Tris, pH 8.8

2mg/ml BSA

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Representative photographs displaying PCR products are shown in Figure 5.

Cloning of PCR Products: In order to further analyze

these PCR products, these products were cloned into a suitable plasmid vector using "TA Cloning" (Invitrogen® The cloning strategy employed here is to directly ligate PCR products into a plasmid vector possessing overhanging T residues at the insertion site, exploiting the fact that Taq polymerase leaves overhanging A residues at the ends of the PCR products. The ligation mixes are transformed into competent E. coli cells and resulting colonies are grown up, plasmid DNA is isolated by the alkaline lysis method (24), and screened by restriction analysis (Figures 6A-6B).

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DNA Sequencing of PCR Products: TA Clones of PCR products were then sequenced by the dideoxy method (25) using Sequenase (U.S. Biochemical). 3-4µg of each plasmid DNA was denatured with NaOH and ethanol precipitated. Labeling reactions were carried out as per the manufacturers recommendations using  $^{35}\text{S-ATP}$ , and the reactions were terminated as per the same protocol. were then analyzed products Sequencing polyacrylamide/7M Urea gels using an IBI sequencing apparatus. Gels were run at 120 watts for 2 hours. Following electrophoresis, the gels were fixed for 15-20 minutes in 10% methanol/10% acetic acid, transferred onto Whatman 3MM paper and dried down in a Biorad® vacuum dryer at 80°C for 2 hours. Gels were then autoradiographed at room temperature for 16-24 hours. In order to determine whether the PCR products were the correct clones, the sequences obtained at the 5' and 3' ends of the molecules were analyzed for the correct 30 primer sequences, as well as adjacent sequences which corresponded to portions of the peptides not used in the design of the primers.

IN-20 was confirmed to be correct and represent a 35 partial cDNA for the PSM gene. In this PCR reaction, I and N primers were used. The DNA sequence reading -55-

from the I primer was:

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ACG GAG CAA AAC TTT CAG CTT GCA AAG (SEQ ID No. K (SEQ ID No. L N

The underlined amino acids were the portion of peptide 6 that was used to design this sense primer and the remaining amino acids which agree with those present within the peptide confirm that this end of the molecule represents the correct protein (PSM antigen).

When analyzed the other end of the molecule by reading from the N primer the anti-sense sequence was:

CTC TTC GGC ATC CCA GCT TGC AAA CAA AAT TGT TCT (SEQ ID 15 No. )

Sense (complementary) Sequence:

- AGA ACA ATT TTG TTT GCA AGC TGG GAT GCC AAG GAG (SEQ ID 20 ) No. A E E (SEQ ID D R T I No. )
- The underlined amino acids here represent the portion 25 of peptide 7 used to create primer N. All of the amino acids upstream of this primer are correct in the IN-20 clone, agreeing with the amino acids found in peptide Further DNA sequencing has enabled us to identify the presence of other PSM peptides within the DNA 30 sequence of the positive clone.

The DNA sequence of this partial cDNA was found to be unique when screened on the Genbank computer database.

CDNA Library Construction and Cloning of Full - Length PSM cDNA: A cDNA library from LNCaP mRNA was constructed using the Superscript® plasmid system The library was transformed using (BRL®-Gibco). competent DH5- $\alpha$  cells and plated onto 100mm plates containing LB plus  $100\mu g/ml$  of Carbenicillin. Plates were grown overnight at 37°C and colonies were transferred to nitrocellulose filters. Filters were processed and screened as per Grunstein and Hogness (26), using the 1.1kb partial cDNA homologous probe which was radiolabelled with 32P-dCTP by random priming (27). Eight positive colonies were obtained which upon DNA restriction and sequencing analysis proved to represent full-length cDNA molecules coding for the PSM antigen. Shown in Figure 7 is an autoradiogram showing the size of the cDNA molecules represented in the library and in Figure 8 restriction analysis of several full-length clones is shown. Figure 9 is a plasmid Southern analysis of the samples in Figure 8, showing that they all hybridize to the 1.1kb partial cDNA probe.

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Both the cDNA as well as the antigen have been screened through the Genbank Computer database (Human Genome Project) and have been found to be unique.

- Northern Analysis of PSM Gene Expression: Northern analysis (28) of the PSM gene has revealed that expression is limited to the prostate and to prostate carcinoma.
- RNA samples (either 10µg of total RNA or 2µg of poly A+RNA) were denatured and electrophoresed through 1.1% agarose/formaldehyde gels at 60 milliamps for 6-8 hours. RNA was then transferred to Nytran® nylon membranes (Schleicher and Schuell®) by pressure blotting in 10x SSC with a Posi-blotter (Stratagene®). RNA was cross-linked to the membranes using a Stratalinker (Stratagene®) and subsequently baked in a

vacuum oven at 80°C for 2 hours. Blots were prehybridized at 65°C for 2 hours in prehybridization solution (BRL®) and subsequently hybridized for 16 hours in hybridization buffer (BRL®) containing 1-2 x 106 cpm/ml of 32 P-labelled random-primed cDNA probe. Membranes were washed twice in 1x SSPE/1% SDS and twice in 0.1x SSPE/1% SDS at 42°C. Membranes were then airdried and autoradiographed for 12-36 hours at -70°C.

PCR Analysis of PSM Gene Expression in Human Prostate
Tissues: PCR was performed on 15 human prostate samples
to determine PSM gene expression. Five samples each
from normal prostate tissue, benign prostatic
hyperplasia, and prostate cancer were used (histology
confirmed by MSKCC Pathology Department).

 $10\mu g$  of total RNA from each sample was reverse transcribed to made cDNA template as previously described in section IV. The primers used corresponded to the 5' and 3' ends of the 1.1kb partial cDNA, IN-20, and therefore the expected size of the amplified band is 1.1kb. Since the  $T_m$  of the primers is 64°C. PCR primers were annealed at 60°C. PCR was carried out for 35 cycles using the same conditions previously described in section IV.

LNCaP and H26 - Ras transfected LNCaP (29) were included as a positive control and DU-145 as a negative control. 14/15 samples clearly amplified the 1.1kb band and therefore express the gene.

#### Experimental Results

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The gene which encodes the 100kD PSM antigen has been identified. The complete cDNA sequence is shown in Sequence ID #1. Underneath that nucleic acid sequence is the predicted translated amino acid sequence. The total number of the amino acids is 750, ID #2. The

hydrophilicity of the predicted protein sequence is shown in Figures 16:1-11. Shown in Figures 17A-17C are three peptides with the highest point of hydrophilicity. They are: Asp-Glu-Leu-Lys-Ala-Glu (SEQ ID No. ); Asn-Glu-Asp-Gly-Asn-Glu (SEQ ID No. ; and Lys-Ser-Pro-Asp-Glu-Gly (SEQ ID No. ).

By the method of Klein, Kanehisa and DeLisi, a specific membrane-spanning domain is identified. The sequence is from the amino acid #19 to amino acid #44: Ala-Gly-Ala-Leu-Val-Leu-Aal-Gly-Gly-Phe-Phe-Leu-Leu-Gly-Phe-Leu-Phe (SEQ ID No. ).

This predicted membrane-spanning domain was computed on PC Gene (computer software program). This data enables prediction of inner and outer membrane domains of the PSM antigen which aids in designing antibodies for uses in targeting and imaging prostate cancer.

When the PSM antigen sequence with other known sequences of the GeneBank were compared, homology between the PSM antigen sequence and the transferrin receptor sequence were found. The data are shown in Figure 18.

Experimental Discussions

Potential Uses for PSM Antigen:

30 1. Tumor detection:

Microscopic:

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Unambiguous tumor designation can be accomplished by use of probes for different antigens. For prostatic cancer, the PSM antigen probe may prove beneficial. Thus PSM could be used for diagnostic purposes and this could be accomplished at the microscopic level using

(control)

in-situ hybridization using sense

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antisense probes derived from the coding region of the cDNA cloned by the applicants. This could be used in assessment of local extraprostatic extension. involvement of lymph node, bone or other metastatic As bone metastasis presents a major problem in prostatic cancer, early detection of metastatic spread is required especially for staging. In some tumors detection of tumor cells in bone marrow portends a grim prognosis and suggests that interventions aimed at metastasis be tried. Detection of PSM antigen expression in bone marrow aspirates or sections may provide such early information. PCR amplification or in-situ hybridization may be used. Using RT-PCR cells in the circulating can be detected by hematogenous metastasis.

2. Antigenic site identification

The knowledge of the cDNA for the antigen also provides for the identification of areas that would serve as good antigens for the development of antibodies for use against specific amino acid sequences of the antigen. Such sequences may be at different regions such as outside, membrane or inside of the PSM antigen. The development of these specific antibodies would provide for immunohistochemical identification of the antigen. These derived antibodies could then be developed for use, especially ones that work in paraffin fixed sections as well as frozen section as they have the greatest utility for immunodiagnosis.

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3. Restriction fragment length polymorphism and genomic DNA

Restriction fragment length polymorphisms (RFLPS) have proven to be useful in documenting the progression of genetic damage that occurs during tumor initiation and promotion. It may be that RFLP analysis will demonstrate that changes in PSM sequence restriction

mapping may provide evidence of predisposition to risk or malignant potential or progression of the prostatic tumor.

Depending on the chromosomal location of the PSM antigen, the PSM antigen gene may serve as a useful chromosome location marker for chromosome analysis.

#### 4. Serum

With the development of antigen specific antibodies, if the antigen or selected antigen fragments appear in the serum they may provide for a serum marker for the presence of metastatic disease and be useful individually or in combination with other prostate specific markers.

#### 5. Imaging

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As the cDNA sequence implies that the antigen has the characteristics of a membrane spanning protein with the majority of the protein on the exofacial surface, antibodies, especially monoclonal antibodies to the peptide fragments exposed and specific to the tumor may provide for tumor imaging local extension of metastatic tumor or residual tumor following prostatectomy or The knowledge of the coding region irradiation. permits the generation of monoclonal antibodies and these can be used in combination to provide for maximal Because the antigen shares a imaging purposes. similarity with the transferrin receptor based on cDNA analysis (approximately 54%), it may be that there is a specific normal ligand for this antigen and that identification of the ligand(s) would provide another means of imaging.

35 6. Isolation of ligands
The PSM antigen can be used to isolate the normal ligand(s) that bind to it. These ligand(s) depending

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on specificity may be used for targeting, or their serum levels may be predictive of disease status. If it is found that the normal ligand for PSM is a carrier molecule then it may be that PSM could be used to bind to that ligand for therapy purposes (like an iron chelating substance) to help remove the ligand from the circulation. If the ligand promotes tumor growth or metastasis then providing soluble PSM antigen would remove the ligand from binding the prostate. Knowledge of PSM antigen structure could lend to generation of small fragment that binds ligand which could serve the same purpose.

#### 7. Therapeutic uses

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The knowledge that the cDNA structure of Ligands. 15 antigen shares structural homology with the transferrin receptor (54% on the nucleic acid level) implies that there may be an endogenous ligand for the receptor that may or may not be transferrin-like. Transferrin is thought to be a ligand that transports 20 iron into the cell after binding to the transferrin receptor. However, apotransferrin is being reported to be a growth factor for some cells which express the transferrin receptor (30). Whether transferrin is a ligand for this antigen or some other ligand binds to 25 this ligand remains to be determined. If a ligand is identified it may carry a specific substance such as a metal ion (iron or zinc or other) into the tumor and thus serve as a means to deliver toxic substances (radioactive or cytotoxic chemical i.e. toxin like 30 ricin or cytotoxic alkylating agent or cytotoxic prodrug) to the tumor.

The main metastatic site for prostatic tumor is the bone. The bone and bone stroma are rich in transferrin. Recent studies suggest that this microenvironment is what provides the right "soil" for

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prostatic metastasis in the bone (31). It may be that this also promotes attachment as well, these factors which reduce this ability may diminish prostatic metastasis to the bone and prostatic metastatic growth in the bone.

It was found that the ligand for the new antigen (thought to be an oncogene and marker of malignant phenotype in breast carcinoma) served to induce differentiation of breast cancer cells and thus could serve as a treatment for rather than promotor of the disease. It may be that ligand binding to the right region of PSM whether with natural ligand or with an antibody may serve a similar function.

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Antibodies against PSM antigen coupled with a cytotoxic agent will be useful to eliminate prostate cancer cells. Transferrin receptor antibodies with toxin conjugates are cytotoxic to a number of tumor cells as tumor cells tend to express increased levels of transferrin receptor (32). Transferrin receptors take up molecules into the cell by endocytosis. Antibody drug combinations can be toxic. Transferrin linked toxin can be toxic.

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Antibodies against PSM antigen coupled with a b) cytotoxic agent will be useful to eliminate prostate The cytotoxic agent may be cancer cells. radioisotope or toxin as known in ordinary skill of the The linkage of the antibody and the toxin or Examples of direct radioisotope can be chemical. linked toxins are doxorubicin, chlorambucil, ricin, pseudomonas exotoxin etc., or a hybrid toxin can be generated % with specificity for PSM and the other % Such a bivalent with specificity for the toxin. molecule can serve to bind to the tumor and the other % to deliver a cytotoxic to the tumor or to bind to and

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activate a cytotoxic lymphocyte such as binding to the T, - T receptor complex. Antibodies of required specificity can also be cloned into T cells and by replacing the immunoglobulin domain of the T cell receptor (TcR); cloning in the desired MAb heavy and light chains; splicing the U, and U, gene segments with the constant regions of the  $\alpha$  and  $\beta$  TCR chains and transfecting these chimeric Ab/TcR genes patients' T cells, propagating these hybrid cells and infusing them into the patient (33). knowledge of tissue specific antigens for targets and generation of MAb's specific for such targets will help make this a usable approach. Because the PSM antigen coding region provides knowledge of the entire coding region, it is possible to generate a number of antibodies which could then be used in combination to achieve an additive or synergistic anti-tumor action. The antibodies can be linked to enzymes which can activate non-toxic prodrugs at its site of the tumor Ab-carboxypeptidase and 4-(bis(2 as chloroethyl) amino) benzoyl-α-glutamic acid and active parent drug in mice (34).

It is possible to produce a toxic genetic chimera such as TP-40 a genetic recombinant that possesses the cDNA from TGF-alpha and the toxic portion of pseudomonas exotoxin so the TGF and portion of the hybrid binds the epidermal growth factor receptor (EGFR) and the pseudomonas portion gets taken up into the cell enzymatically and inactivates the ribosomes ability to perform protein synthesis resulting in cell death.

In addition, once the ligand for the PSM antigen is identified, toxin can be chemically conjugated to the ligands. Such conjugated ligands can be therapeutically useful. Examples of the toxins are daunomycin, chlorambucil, ricin, pseudomonas exotoxin,

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etc. Alternatively, chimeric construct can be created linking the cDNA of the ligand with the cDNA of the toxin. An example of such toxin is  $TGF\alpha$  and pseudomonas exotoxin (35).

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#### 8. Others

The PSM antigen may have other uses. It is well known that the prostate is rich in zinc, if the antigen provides function relative to this or other biologic function the PSM antigen may provide for utility in the treatment of other prostatic pathologies such as benign hyperplastic growth and/or prostatitis.

Because purified PSM antigen can be generated, the purified PSM antigen can be linked to beads and use it 15 like a standard "affinity" purification. Serum, urine or other biological samples can be used to incubate with the PSM antigen bound onto beads. The beads may be washed thoroughly and then eluted with salt or pH gradient. The eluted material is SDS gel purified and 20 used as a sample for microsequencing. The sequences will be compared with other known proteins and if unique, the technique of degenerated PCR can be employed for obtaining the ligand. Once known, the affinity of the ligand will be determined by standard 25 protocols (15).

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#### EXAMPLE 2:

# EXPRESSION OF THE PROSTATE SPECIFIC MEMBRANE ANTIGEN

A 2.65 kb complementary DNA encoding PSM was cloned. 5 Immunohistochemical analysis of the LNCaP, DU-145, and PC-3 prostate cancer cell lines for PSM expression using the 7E11-C5.3 antibody reveals intense staining in the LNCaP cells, with no detectable expression in Coupled in-vitro both the DU-145 and PC-3 cells. 10 transcription/ translation of the 2.65 kb full-length PSM cDNA yields an 84 kDa protein corresponding to the predicted polypeptide molecular weight of PSM. Postprotein with translational modification of this pancreatic canine microsomes yields the expected 100 15 kDa PSM antigen. Following transfection of PC-3 cells eukaryotic . with the full-length PSM cDNA in a expression vector applicant's detect expression of the PSM glycoprotein by Western analysis using the 7Ell-C5.3 monoclonal antibody. Ribonuclease protection 20 analysis demonstrates that the expression of PSM mRNA is almost entirely prostate-specific in human tissues. PSM expression appears to be highest in hormonedeprived states and is hormonally modulated steroids, with DHT downregulating PSM expression in the 25 human prostate cancer cell line LNCaP by 8-10 fold, testosterone downregulating PSM by 3-4 fold, and corticosteroids showing no significant effect. Normal and malignant prostatic tissues consistently show high PSM expression, whereas heterogeneous, and at times 30 absent, from expression of PSM in benign prostatic hyperplasia. LNCaP tumors implanted and grown both orthotopically and subcutaneously in nude mice, abundantly express PSM providing an excellent in-vivo model system to study the regulation and modulation of 35 PSM expression.

## Materials and Methods:

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Cells and Reagents: The LNCaP, DU-145, and PC-3 cell lines were obtained from the American Type Culture Details regarding the establishment and Collection. these cell lines have characteristics of previously published (5A,7A,8A). Unless specified otherwise, LNCaP cells were grown in RPMI 1640 media supplemented with L-glutamine, nonessential 5% fetal calf serum (Gibco-BRL, and Gaithersburg, MD.) in a CO, incubator at 37C. DU-145 and PC-3 cells were grown in minimal essential medium supplemented with 10% fetal calf serum. All cell media were obtained from the MSKCC Media Preparation Restriction and modifying enzymes were purchased from Gibco-BRL unless otherwise specified.

Immunohistochemical Detection of PSM: Avidin-biotin method of detection was employed to analyze prostate cancer cell lines for PSM antigen expression (9A). Cell cytospins were made on glass slides using 5x104 cells/100ul per slide. Slides were washed twice with PBS and then incubated with the appropriate suppressor serum for 20 minutes. The suppressor serum was drained off and the cells were incubated with diluted 7E11-C5.3 (5g/ml) monoclonal antibody for 1 hour. Samples were then washed with PBS and sequentially incubated with secondary antibodies for 30 minutes and with avidinbiotin complexes for 30 minutes. Diaminobenzidine served as the chromogen and color development followed by hematoxylin counterstaining and mounting. Duplicate controls cytospins were used as for a positive control, the experiment. As cytokeratin monoclonal antibody CAM 5.2 was used following the same procedure described above. Human EJ bladder carcinoma cells served as a negative control.

In-Vitro Transcription/Translation of PSM Antigen: Plasmid 55A containing the full length 2.65 kb PSM cDNA in the plasmid pSPORT 1 (Gibco-BRL) was transcribed invitro using the Promega TNT system (Promega Corp. Madison, WI). T7 RNA polymerase was added to the cDNA in a reaction mixture containing rabbit reticulocyte lysate, an amino acid mixture lacking methionine, buffer, and 35S-Methionine (Amersham) and incubated at 30C for 90 minutes. Post-translational modification of the resulting protein was accomplished by the addition of pancreatic canine microsomes into the reaction mixture (Promega Corp. Madison, WI.). Protein products were analyzed by electrophoresis on 10% SDS-PAGE gels subsequently treated with were which autoradiography enhancer (Amersham, Arlington Heights, IL.) according to the manufacturers instructions and in a vacuum dryer. Gels were 80C autoradiographed overnight at -70C using Hyperfilm MP (Amersham).

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Transfection of PSM into PC-3 Cells: The full length PSM cDNA was subcloned into the pREP7 eukaryotic expression vector (Invitrogen, San Diego, Plasmid DNA was purified from transformed DH5-alpha bacteria (Gibco-BRL) using Qiagen maxi-prep plasmid isolation columns (Qiagen Inc., Chatsworth, CA.). Purified plasmid DNA (6-10g) was diluted with 900ul of Optimem media (Gibco-BRL) and mixed with 30ul of which had (Gibco-BRL) reagent Lipofectin previously diluted with 900l of Optimem media. mixture was added to T-75 flasks of 40-50% confluent PC-3 cells in Optimem media. After 24-36 hours, cells into 100mm dishes trypsinized and split containing RPMI 1640 media supplemented with 10% fetal calf serum and 1 mg/ml of Hygromycin B (Calbiochem, La The dose of Hygromycin B used was Jolla, CA.). previously determined by a time course/dose response

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cytotoxicity assay. Cells were maintained in this media for 2-3 weeks with changes of media and Hygromycin B every 4-5 days until discrete colonies appeared. Colonies were isolated using 6mm cloning cylinders and expanded in the same media. As a control, PC-3 cells were also transfected with the pREP7 plasmid alone. RNA was isolated from the transfected cells and PSM mRNA expression was detected by both RNase Protection analysis (described later) and by Northern analysis.

Western Blot Detection of PSM Expression: Crude protein lysates were isolated from LNCaP, PC-3, and PSMtransfected PC-3 cells as previously described (10A). LNCaP cell membranes were also isolated according to 15 published methods (10A). Protein concentrations were quantitated by the Bradford method using the BioRad protein reagent kit (BioRad, Richmond, CA.). Following denaturation,  $20\mu g$  of protein was electrophoresed on a 20 10% SDS-PAGE gel at 25 mA for 4 hours. electroblotted onto Immobilon P membranes (Millipore, Bedford, MA.) overnight at 4C. Membranes were blocked in 0.15M NaCl/0.01M Tris-HCl (TS) plus 5% BSA followed by a 1 hour incubation with 7E11-C5.3 monoclonal antibody  $(10\mu g/ml)$ . 25 Blots were washed 4 times with 0.15M NaCl/0.01M Tris-HCl/0.05% Triton-X 100 (TS-X) and incubated for 1 hour with rabbit anti-mouse IqG (Accurate Scientific, Westbury, N.Y.) at concentration of  $10\mu g/ml$ .

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Blots were then washed 4 times with TS-X and labeled with <sup>125</sup>I-Protein A (Amersham, Arlington Heights, IL.) at a concentration of 1 million cpm/ml. Blots were then washed 4 times with TS-X and dried on Whatman 3MM paper, followed by overnight autoradiography at -70C using Hyperfilm MP (Amersham).

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Orthotopic and Subcutaneous LNCaP Tumor Growth in Nude Mice: LNCaP cells were harvested from sub-confluent cultures by a one minute exposure to a solution of 0.25% trypsin and 0.02% EDTA. Cells were resuspended in RPMI 1640 media with 5% fetal bovine serum, washed (Collaborative and diluted in either Matrigel Biomedical Products, Bedford, MA.) or calcium and magnesium-free Hank's balanced salt solution (HBSS). Only single cell suspensions with greater than 90% viability by trypan blue exclusion were used for in vivo injection. Male athymic Swiss (nu/nu) nude mice 4-6 weeks of age were obtained from the Memorial Sloan-Kettering Cancer Center Animal Facility. subcutaneous tumor cell injection one million LNCaP cells resuspended in 0.2 mls. of Matrigel were injected into the hindlimb of each mouse using a disposable syringe fitted with a 28 gauge needle. For orthotopic injection, mice were first anesthetized with an intraperitoneal injection of Pentobarbital and placed in the supine position. The abdomen was cleansed with Betadine and the prostate was exposed through a midline 2.5 million LNCaP tumor cells in 0.1 ml. incision. were injected directly into either posterior lobe using a 1 ml disposable syringe and a 28 gauge needle. LNCaP cells with and without Matrigel were injected. 25 Abdominal closure was achieved in one layer using Autoclip wound clips (Clay Adams, Parsippany, N.J.). were harvested in 6-8 weeks, confirmed histologically by faculty of the Memorial Sloan-Kettering Cancer Center Pathology Department, 30 frozen in liquid nitrogen for subsequent RNA isolation.

RNA Isolation: Total cellular RNA was isolated from cells and tissues by standard techniques (11,12) as well as by using RNAzol B (Cinna/Biotecx, Houston, TX.). RNA concentrations and quality were assessed by UV spectroscopy on a Beckman DU 640 spectrophotometer WO 96/26272 PCT/US96/02424

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and by gel analysis. Human tissue total RNA samples were purchased from Clontech Laboratories, Inc., Palo Alto, CA.

5 Ribonuclease Protection Assays: A portion of the PSM cDNA was subcloned into the plasmid vector pSPORT 1 (Gibco-BRL) and the orientation of the cDNA insert relative to the flanking T7 and SP6 RNA polymerase promoters was verified by restriction analysis. Linearization of this plasmid upstream of the PSM 10 insert followed by transcription with polymerase yields a 400 nucleotide antisense RNA probe, of which 350 nucleotides should be protected from RNase digestion by PSM RNA. This probe was used in Figure Plasmid IN-20, containing a 1 kb partial PSM cDNA 15 in the plasmid pCR II (Invitrogen) was also used for riboprobe synthesis. IN-20 linearized with Xmn I (Gibco-BRL) yields a 298 nucleotide anti-sense RNA probe when transcribed using SP6 RNA polymerase, of 20 which 260 nucleotides should be protected from RNase digestion by PSM mRNA. This probe was used in Figures 21 and 22. Probes were synthesized using SP6 RNA polymerase (Gibco-BRL), rNTPs (Gibco-BRL), (Promega), and 32P-rCTP (NEN, Wilmington, DE.) according 25 to published protocols (13). Probes were purified over NENSORB 20 purification columns (NEN) and approximately 1 million cpm of purified, radiolabeled PSM probe was mixed with  $10\mu$  of each RNA and hybridized overnight at 45C using buffers and reagents from the RPA II kit (Ambion, Austin, TX). 30 Samples were processed as per manufacturer's instructions and analyzed on 5% polyacrilamide/7M urea denaturing gels using Seg ACRYL reagents (ISS, Natick, MA.). Gels were pre-heated to 55C and run for approximately 1-2 hours at 25 watts. Gels were then fixed for 30 minutes in 10% methanol/10% 35 acetic acid, dried onto Whatman 3MM paper at 80C in a BioRad vacuum dryer and autoradiographed overnight with

Hyperfilm MP (Amersham). Quantitation of PSM expression was determined by using a scanning laser densitometer (LKB, Piscataway, NJ.).

Steroid Modulation Experiment: LNCaP cells (2 million) 5 were plated onto T-75 flasks in RPMI 1640 media supplemented with 5% fetal calf serum and grown 24 hours until approximately 30-40% confluent. were then washed several times with phophate-buffered saline and RPMI medium supplemented with 5% charcoal-10 extracted serum was added. Cells were then grown for another 24 hours, at which time dihydrotesterone, progesterone, estradiol, testosterone, dexamethasone (Steraloids Inc., Wilton, NH.) were added at a final concentration of 2 nM. Cells were grown for 15 another 24 hours and RNA was then harvested as previously described and PSM expression analyzed by ribonuclease protection analysis:

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## Experimental Results

Immunohistochemical Detection of PSM: Using the 7Ell-C5.3 anti-PSM monoclonal antibody, PSM expression is clearly detectable in the LNCaP prostate cancer cell line, but not in the PC-3 and DU-145 cell lines (Figures 17A-17C). All normal and malignant prostatic tissues analyzed stained positively for PSM expression.

In-Vitro Transcription/Translation of PSM Antigen: As shown in Figure 18, coupled in-vitro transcription/ translation of the 2.65 kb full-length PSM cDNA yields an 84 kDa protein species in agreement with the expected protein product from the 750 amino acid PSM open reading frame. Following post-translational modification using pancreatic canine microsomes were obtained a 100 kDa glycosylated protein species

consistent with the mature, native PSM antigen.

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Transfected PC-3 Cells: PC-3 cells transfected with the full length PSM cDNA in the pREP7 expression vector were assayed for expression of SM mRNA by Northern analysis. A clone with high PSM mRNA expression was selected for PSM antigen analysis by Western blotting using the 7E11-C5.3 antibody. In Figure 19, the 100 kDa PSM antigen is well expressed in LNCaP cell lysate and membrane fractions, as well as in PSM-transfected PC-3 cells but not in native PC-3 cells. This detectable expression in the transfected PC-3 cells proves that the previously cloned 2.65 kb PSM cDNA encodes the antigen recognized by the 7E11-C5.3 antiprostate monoclonal antibody.

PSM mRNA Expression: Expression of PSM mRNA in normal analyzed using ribonuclease tissues was Tissue expression of PSM appears protection assays. predominantly within the prostate, with very low levels of expression detectable in human brain and salivary gland (Figure 20). No detectable PSM mRNA expression evident in non-prostatic human tissues when analyzed by Northern analysis. On occasion it is noted that detectable PSM expression in normal human small intestine tissue, however this mRNA expression is variable depending upon the specific riboprobe used. samples of normal human prostate and human prostatic adenocarcinoma assayed have revealed clearly detectable PSM expression, whereas generally decreased or absent expression of PSM in tissues exhibiting benign hyperplasia (Figure 21). In human LNCaP tumors grown both orthotopically and subcutaneously in nude mice abundant PSM expression with or without the use of matrigel, which is required for the growth of subcutaneously implanted LNCaP cells was detected

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PSM mRNA expression is distinctly (Figure 21). modulated by the presence of steroids in physiologic doses (Figure 22). DHT downregulated expression by 8-10 fold after 24 hours and testosterone diminished PSM Estradiol and progesterone expression by 3-4 fold. also downregulated PSM expression in LNCaP cells, perhaps as a result of binding to the mutated androgen receptor known to exist in the LNCaP cell. Overall, PSM expression is highest in the untreated LNCaP cells grown in steroid-depleted media, a situation that simulates the hormone-deprived (castrate) state in-This experiment was repeated at steroid dosages ranging from 2-200 nM and at time points from 6 hours to 7 days with similar results; maximal downregulation of PSM mRNA was seen with DHT at 24 hours at doses of 2-20 nM.

# Experimental Discussion

Previous research has provided two valuable prostatic bio-markers, PAP and PSA, both of which have had a significant impact on the diagnosis, treatment, and management of prostate malignancies. The present work describing the preliminary characterization of the prostate-specific membrane antigen (PSM) reveals it to be a gene with many interesting features. almost entirely prostate-specific as are PAP and PSA, and as such may enable further delineation of the unique functions and behavior of the prostate. The predicted sequence of the PSM protein (3) and its presence in the LNCaP cell membrane as determined by Western blotting and immunohistochemistry, indicate that it is an integral membrane protein. Thus, PSM provides an attractive cell surface epitope for antibody-directed diagnostic imaging and cytotoxic targeting modalities (14). The ability to synthesize the PSM antigen in-vitro and to produce

xenografts maintaining high levels of PSM expression provides us with a convenient and attractive model system to further study and characterize the regulation and modulation of PSM expression. Also, the high level of PSM expression in the LNCaP cells provides an excellent in-vitro model system. Since PSM expression is hormonally-responsive to steroids and may be highly expressed in hormone-refractory disease (15). detection of PSM mRNA expression in minute quantities in brain, salivary gland, and small intestine warrants further investigation, although these tissues were expression PSM antigen of for immunohistochemistry using the 7E11-C5.3 antibody (16). In all of these tissues, particularly small intestine, mRNA expression using a probe corresponding to a region of the PSM cDNA near the 3' end, whereas expression when using a 5' end PSM probe was not detected. results may indicate that the PSM mRNA transcript undergoes alternative splicing in different tissues.

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Applicants approach is based on prostate tissue enzyme or cytokine chimeras. specific promotor: Promotor specific activation of prodrugs such as non toxic gancyclovir which is converted to a toxic metabolite by herpes simplex thymidine kinase or the prodrug 4-(bis(2chloroethyl)amino)benzoyl-1-glutamic acid to the benzoic acid mustard alkylating agent by the pseudomonas carboxy peptidase G2 was examined. these drugs are activated by the enzyme (chimera) specifically in the tumor the active drug is released only locally in the tumor environment, destroying the surrounding tumor cells. Promotor specific activation IL-2 or GM-CSF for of cytokines such as IL-12, activation and specific antitumor vaccination is Lastly the tissue specific promotor examined. activation of cellular death genes may also prove to be useful in this area.

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Gene Therapy Chimeras: The establishment of "chimeric DNA" for gene therapy requires the joining of different segments of DNA together to make a new DNA that has characteristics of both precursor DNA species involved in the linkage. In this proposal the two pieces being linked involve different functional aspects of DNA, the promotor region which allows for the reading of the DNA for the formation of mRNA will provide specificity and the DNA sequence coding for the mRNA will provide for therapeutic functional DNA.

DNA-Specified Enzyme or Cytokine mRNA: When effective, antitumor drugs can cause the regression of very large amounts of tumor. The main requirements for antitumor drug activity is the requirement to achieve both a long enough time (t) and high enough concentration (c) (cxt) of exposure of the tumor to the toxic drug to assure sufficient cell damage for cell death to occur. drug also must be "active" and the toxicity for the tumor greater than for the hosts normal cells (22). The availability of the drug to the tumor depends on tumor blood flow and the drugs diffusion ability. does not provide for. Blood flow to the tumor selectivity as blood flow to many normal tissues is often as great or greater than that to the tumor. majority of chemotherapeutic cytotoxic drugs are often as toxic to normal tissue as to tumor tissue. Dividing cells are often more sensitive than non-dividing normal cells, but in many slow growing solid tumors such as prostatic cancer this does not provide for antitumor specificity (22).

Previously a means to increase tumor specificity of antitumor drugs was to utilize tumor associated enzymes to activate nontoxic prodrugs to cytotoxic agents (19). A problem with this approach was that most of the enzymes found in tumors were not totally specific in

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their activity and similar substrate active enzymes or the same enzyme at only slightly lower amounts was found in other tissue and thus normal tissues were still at risk for damage.

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To provide absolute specificity and unique activity. viral, bacterial and fungal enzymes which have unique specificity for selected prodrugs were found which were not present in human or other animal cells. Attempts to utilize enzymes such as herpes simplex thymidine bacterial cytosine deaminase carboxypeptidase G-2 were linked to antibody targeting systems with modest success (19). Unfortunately, antibody targeted enzymes limit the number of enzymes available per cell. Also, most antibodies do not have a high tumor target to normal tissue ratio thus normal tissues are still exposed reducing the specificity of these unique enzymes. Antibodies are large molecules that have poor diffusion properties and the addition of the enzymes molecular weight further reduces the antibodies diffusion.

Gene therapy could produce the best desired result if it could achieve the specific expression of a protein in the tumor and not normal tissue in order that a high local concentration of the enzyme be available for the production in the tumor environment of active drug (21).

#### 30 Cytokines:

Results demonstrated that tumors such as the bladder and prostate were not immunogenic, that is the administration of irradiated tumor cells to the animal prior to subsequent administration of non-irradiated tumor cells did not result in a reduction of either the number of tumor cells to produce a tumor nor did it reduce the growth rate of the tumor. But if the tumor

was transfected with a retrovirus and secreted large concentrations of cytokines such as Il-2 then this could act as an antitumor vaccine and could also reduce the growth potential of an already established and IL-2 was the best, GM-CSF also had growing tumor. 5 activity whereas a number of other cytokines were much less active. In clinical studies just using IL-2 for immunostimulation, very large concentrations had to be given which proved to be toxic. The key to the success of the cytokine gene modified tumor cell is that the 10 cytokine is produced at the tumor site locally and is not toxic and that it stimulates immune recognition of the tumor and allows specific and non toxic recognition and destruction of the tumor. The exact mechanisms of how IL-2 production by the tumor cell activates immune 15 fully understood, not recognition is explanation is that it bypasses the need for cytokine production by helper T cells and directly stimulates CD8 cytotoxic activated antigen tumor Activation of antigen presenting cells may also occur. 20

# Tissue Promotor-Specific Chimera DNA Activation

# Non-Prostatic Tumor Systems:

It has been observed in non-prostatic tumors that the 25 use of promotor specific activation can selectively lead to tissue specific gene expression of In melanoma the use of transfected gene. tyrosinase promotor which codes for responsible for melanin expression produced over a 50 30 fold greater expression of the promotor driven reporter gene expression in melanoma cells and not non melanoma Similar specific activation was seen in the melanoma cells transfected when they were growing in mice. In that experiment no non-melanoma or melanocyte 35 cell expressed the tyrosinase drive reporter gene The research group at Welcome Laboratories

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have cloned and sequenced the promoter region of the gene coding for carcinoembryonic antigen (CEA). CEA is expressed on colon and colon carcinoma cells but specifically on metastatic. A gene chimera was generated which cytosine deaminase. Cytosine deaminase which converts 5 flurorocytosine into 5 fluorouracil and observed a large increase in the ability to selectively kill CEA promotor driven colon tumor cells but not normal liver cells. In vivo they observed that bystander tumor cells which were not transfected with the cytosine deaminase gene were also killed, and that there was no toxicity to the host animal as the large tumors were regressing following treatment. (HSV), thymidine kinase similarly simplex virus, activates the prodrug gancyclovir to be toxic towards dividing cancer cells and HSV thymidine kinase has been shown to be specifically activatable by tissue specific promoters.

Prostatic Tumor Systems: The therapeutic key to 20 effective cancer therapy is to achieve specificity and spare the patient toxicity. Gene therapy may provide a key part to specificity in that non-essential tissues such as the prostate and prostatic tumors produce 25 tissue specific proteins, such as acid phosphatase (PAP), prostate specific antigen (PSA), and a gene which was cloned, prostate-specific membrane antigen (PSM). Tissues such as the prostate contain selected specific transcription factors which responsible for binding to the promoter region of the 30 DNA of these tissue specific mRNA. The promoter for PSA has been cloned. Usually patients who are being treated for metastatic prostatic cancer have been put on androgen deprivation therapy which dramatically reduces the expression of mRNA for PSA. PSM on the 35 other hand increases in expression with hormone deprivation which-means it would be even more intensely WO 96/26272

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expressed on patients being treated with hormone therapy.

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#### EXAMPLE 3:

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# Sensitive Detection of Prostatic Hematogenous Micrometastases Using PSA and PSM-Derived Primers in the Polymerase Chain Reaction

A PCR-based assay was developed enabling sensitive detection of hematogenous micrometastases in patients with prostate cancer. "Nested PCR", was performed by amplifying mRNA sequences unique to prostate-specific antigen and to the prostate-specific membrane antigen, and compared have their respective results. Micrometastases were detected in 2/30 patients (6.7%) by PCR with PSA-derived primers, while PSM-derived primers detected tumor cells in 19/16 patients (63.3%). All 8 negative controls were negative with both PSA and PSM PCR. Assays were repeated to confirm results, and PCR products were verified by DNA sequencing and Southern analysis. Patients harboring circulating prostatic tumor cells as detected by PSM, and not by PSA-PCR included 4 patients previously treated with radical prostatectomy and with non-measurable serum PSA levels at the time of this assay. The significance of findings with respect to future recurrence and progression will be investigated.

Improvement in the overall survival of patients with prostate cancer will depend upon earlier diagnosis. Localized disease, without evidence of extra-prostatic spread, is successfully treated with either radical prostatectomy or external beam radiation, with excellent long-term results (2,3). The major problem is that approximately two-thirds of men diagnosed with prostate cancer already have evidence of advanced extra-prostatic spread at the time of diagnosis, for which there is at present no cure (4). The use of clinical serum markers such as prostate-specific

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antigen (PSA) and prostatic acid phosphatase (PAP) have enabled clinicians to detect prostatic carcinomas earlier and provide useful parameters to follow responses to therapy (5). Yet, despite the advent of sensitive serum PSA assays, radionuclide bone scans, CT scans and other imaging modalities, results have not detected the presence of micrometastatic cells prior to their establishment of solid metastases. Previous work has been done utilizing the polymerase chain reaction to amplify mRNA sequences unique to breast, leukemia, and other malignant cells in the circulation and enable early detection of micrometastases (6,7). Recently, a PCR-based approach utilizing primers derived from the PSA DNA sequence was published (8). In this study 3/12 patients with advanced, stage D prostate cancer had detectable hematogenous micrometastases.

PSM appears to be an integral membrane glycoprotein which is very highly expressed in prostatic tumors and metastases and is almost entirely prostate-specific (10). Many anaplastic tumors and bone metastases have variable and at times no detectable expression of PSA, whereas these lesions appear to consistently express Prostatic tumor cells that escape high levels of PSM. from the prostate gland and enter the circulation are likely to have the potential to form metastases and are possibly the more aggressive and possibly anaplastic cells, a population of cells that may not express high levels of PSA, but may retain high expression of PSM. DNA primers derived from the sequences of both PSA and PSM in a PCR assay were used to detect micrometastatic cells in the peripheral circulation. Despite the high level of amplification and sensitivity of conventional RNA PCR, "Nested" PCR approach in which a amplified target sequence was employed, and subsequently use this PCR product as the template for another round of PCR amplification with a new set of primers totally WO 96/26272 PCT/US96/02424

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contained within the sequence of the previous product. This approach has enabled us to increase the level of detection from one prostatic tumor cell per 10,000 cells to better than one cell per ten million cells.

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#### Materials and Methods

Cells and Reagents: LNCaP and MCF-7 cells were obtained from the American Type Culture Collection (Rockville, establishment Details regarding the MD.). these cell lines have been characteristics of previously published (11,12). Cells were grown in RPMI 1640 media supplemented with L-glutamine, nonessential amino acids, obtained from the MSKCC Media Preparation Facility, and 5% fetal calf (Gibco-BRL, serum Gaithersburg, MD.) in a CO, incubator at 37C. All cell media was obtained from the MSKCC Media Preparation Routine chemical reagents were of the highest grade possible and were obtained from Sigma Chemical Company, St. Louis, MO.

Patient Blood Specimens: All blood specimens used in this study were from patients seen in the outpatient offices of urologists on staff at MSKCC. coagulated (purple top) tubes per patient were obtained at the time of their regularly scheduled blood draws. Specimen procurement was conducted as per the approval of the MSKCC Institutional Review Board. Samples were promptly brought to the laboratory for immediate Serum PSA and PAP determinations were processing. performed by standard techniques by the MSKCC Clinical determinations Chemistry Laboratory. PSA performed using the Tandem PSA assay (Hybritech, San The eight blood specimens used as Diego, CA.). negative controls were from 2 males with normal serum PSA values and biopsy-proven BPH, one healthy female, 3 healthy males, one patient with bladder cancer, and one patient with acute promyelocytic leukemia.

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Blood Sample Processing/RNA Extraction: 4 ml of whole anticoagulated venous blood was mixed with 3 ml of ice cold phosphate buffered saline and then carefully layered atop 8 ml of Ficoll (Pharmacia, Uppsala, Sweden) in a 15-ml polystyrene tube. Tubes were centrifuged at 200 x g for 30 min. at 4C. sterile pasteur pipette, the buffy coat layer (approx. 1 ml.) was carefully removed and rediluted up to 50 ml with ice cold phosphate buffered saline in a 50 ml polypropylene tube. This tube was then centrifuged at for 30 min at 4C. The supernatant was 2000 x q carefully decanted and the pellet was allowed to drip One ml of RNazol B was then added to the pellet and total RNA was isolated as per manufacturers directions (Cinna/Biotecx, Houston, TX.). RNA. concentrations and purity were determined by UV spectroscopy on a Beckman DU 640 spectrophotometer and by gel analysis.

Determination of PCR Sensitivity: RNA was isolated from LNCaP cells and from mixtures of LNCaP and MCF-7 cells at fixed ratios (i.e. 1:100, 1:1000, etc.) using RNAzol B. Nested PCR was then performed as described below with both PSA and PSM primers in order to determine the limit of detection for the assay. LNCaP:MCF-7 (1:100,000) cDNA was diluted with distilled water to obtain concentrations of 1:1,000,000 and 1:10,000,000. MCF-7 cells were chosen because they have been previously tested and shown not to express PSM by PCR.

polymerase Chain Reaction: The PSA outer primers used span portions of exons 4 and 5 to yield a 486 bp PCR product and enable differentiation between cDNA and possible contaminating genomic DNA amplification. The upstream primer sequence beginning at nucleotide 494 in

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PSA cDNA sequence is 5'-TACCCACTGCATCAGGAACA-3' (SEQ. ID. No. ) and the downstream primer at nucleotide 960 is 5'-CCTTGAAGCACCACTTACA-3' (SEQ. ID. No. ). PSA inner upstream primer (beginning at nucleotide 559) 5'-ACACAGGCCAGGTATTTCAG-3' (SEQ. ID. No. ) and the nucleotide 894) downstream primer (at GTCCAGCGTCCAGCACACAG-3' (SEQ. ID. No. ) yield a 355 bp PCR product. All primers were synthesized by the MSKCC Microchemistry Core Facility.  $5\mu q$  of total RNA was reverse-transcribed into cDNA in a total volume of 20µl using Superscript reverse transcriptase (Gibco-BRL) according to the manufacturers recommendations. this cDNA served as the starting template for the outer primer PCR reaction. The 20µl PCR mix included: 0.5U Taq polymerase (Promega Corp., Madison, WI.), Promega reaction buffer, 1.5mM MgCl<sub>2</sub>, 200mM dNTPs, and 1.0μM of each primer. This mix was then transferred to a Perkin Elmer 9600 DNA thermal cycler and incubated for 25 The PCR profile was as follows: 94C x 15 sec., 60C x 15 sec., and 72C for 45 sec. cycles, samples were placed on ice, and 1µl of this reaction mix served as the template for another round of PCR using the inner primers. The first set of tubes were returned to the thermal cycler for 25 additional cycles. PSM-PCR required the selection of primer pairs that also spanned an intron in order to be certain that cDNA and not genomic DNA were being amplified.

The PSM outer primers yield a 946 bp product and the inner primers a 434 bp product. The PSM outer upstream primer used was 5'-ATGGGTGTTTGGTGGTATTGACC-3' (SEQ. ID. No. ) (beginning at nucleotide 1401) and the downstream 2348) 5'-(at nucleotide was primer TGCTTGGAGCATAGATGACATGC-3' (SEQ. ID. No. ) The PSM inner upstream primer (at nucleotide 1581) was 5'-ACTCCTTCAAGAGCGTGGCG-3' (SEQ. ID. No. and the (at nucleotide 2015) 5'downstream primer was

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AACACCATCCCTCCTCGAACC-3'(SEQ. ID. No. ). was the same as for the PSA assay. The 501 PCR mix included: 1U Taq Polymerase (Promega), 250M dNTPs, 10mM -mercaptoethanol, 2mM MgCl,, and 5l of a 10x buffer mix containing: 166mM NH,SO,, 670mM Tris pH 8.8, and 2 mg/ml of acetylated BSA. PCR was carried out in a Perkin Elmer 480 DNA thermal cycler with the following parameters: 94C x 4 minutes for 1 cycle, 94C x 30 sec., 58C  $\times$  1 minute, and 72C  $\times$  1 minute for 25 cycles, followed by 72C x 10 minutes. Samples were then iced and 21 of this reaction mix was used as the template another 25 cycles with a new reaction mix containing the inner PSM primers. cDNA quality was verified by performing control reactions using primers derived from -actin yielding a 446 bp PCR product. upstream primer used was 5'-AGGCCAACCGCGAGAAGATGA-3' (SEQ. ID. No. ) (exon 3) and the downstream primer was 5'-ATGTCACACTGGGGAAGC-3' (SEQ. ID. No. ) (exon 4). The entire PSA mix and 101 of each PSM reaction mix were run on 1.5-2% agarose gels, stained with ethidium bromide and photographed in an Eagle Eye Video Imaging System (Stratagene, Torrey Pines, CA.). Assays were repeated at least 3 times to verify results.

Cloning and Sequencing of PCR Products: PCR products 25 were cloned into the pCR II plasmid vector using the TA cloning system (Invitrogen). These plasmids were transformed into competent E. coli cells using standard methods (13) and plasmid DNA was isolated using Magic and screened by restriction Minipreps (Promega) 30 analysis. TA clones were then sequenced by the dideoxy method (14) using Sequenase (U.S. Biochemical). of each plasmid was denatured with NaOH and ethanol Labeling reactions were carried out precipitated. according to the manufacturers recommendations using 35  $^{35}\text{S-dATP}$  (NEN), and the reactions were terminated as discussed in the same protocol. Sequencing products were then analyzed on 6% polyacrilamide/7M urea gels run at 120 watts for 2 hours. Gels were fixed for 20 minutes in 10% methanol/10% acetic acid, transferred to Whatman 3MM paper and dried down in a vacuum dryer for 2 hours at 80C. Gels were then autoradiographed at room temperature for 18 hours.

Southern Analysis: Ethidium-stained agarose gels of PCR products were soaked for 15 minutes in 0.2N HCl, followed by 30 minutes each in 0.5N NaOH/1.5M NaCl and Gels were then Tris pH 7.5/1.5M NaCl. equilibrated for 10 minutes in 10x SSC (1.5M NaCl/0.15M Sodium Citrate. DNA was transferred onto Nytran nylon (Schleicher and Schuell) by pressure membranes blotting in 10x SSC with a Posi-blotter (Stratagene). DNA was cross-linked to the membrane using a UV Blots were pre-hybridized Stratalinker (Stratagene). at 65C for 2 hourthes and subsequently hybridized with denatured 32P-labeled, random-primed cDNA probes (either Blots were washed twice in lx PSM or PSA) (9,15). SSPE/0.5% SDS at 42C and twice in 0.1x SSPE/0.5% SDS at 50C for 20 minutes each. Membranes were air-dried and autoradiographed for 30 minutes to 1 hour at -70C with Kodak X-Omat film.

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### Experimental Results

PCR amplification with nested primers improved the level of detection of prostatic cells from approximately one prostatic cell per 10,000 MCF-7 cells to better than one cell per million MCF-7 cells, using either PSA or PSM-derived primers (Figures 26 and 27). This represents a substantial improvement in the ability to detect minimal disease. Characteristics of the 16 patients analyzed with respect to their clinical stage, treatment, serum PSA and PAP values, and results of the assay are shown. In total, PSA-PCR detected

tumor cells in 2/30 patients (6.7%), whereas PSM-PCR detected cells in 19/30 patients (63.3%). There were no patients positive for tumor cells by PSA and not by PSM, while PSM provided 8 positive patients not detected by PSA. Patients 10 and 11 in table 1, both 5 with very advanced hormone-refractory disease were detected by both PSA and PSM. Both of these patients have died since the time these samples were obtained. .Patients 4, 7, and 12, all of whom were treated with radical prostatectomies for clinically localized 10 disease, and all of whom have non-measurable serum PSA values 1-2 years postoperatively were positive for circulating prostatic tumor cells by PSM-PCR, negative by PSA-PCR. A representative ethidium stained gel photograph for PSM-PCR is shown in Figure 28. 15 Samples run in lane A represent PCR products generated from the outer primers and samples in lanes labeled B are products of inner primer pairs. The corresponding PSM Southern blot autoradiograph is shown in Figure 29. The sensitivity of the Southern blot analysis exceeded 20 that of ethidium staining, as can be seen in several samples where the outer product is not visible on Figure 28, but is detectable by Southern blotting as shown in Figure 29. In addition, sample 3 on Figures 28 and 29 (patient 6 in Figure 30) appears to contain 25 both outer and inner bands that are smaller than the corresponding bands in the other patients. sequencing has confirmed that the nucleotide sequence of these bands matches that of PSM, with the exception This may represent either an of a small deletion. 30 artifact of PCR, alternative splicing of PSM mRNA in this patient, or a PSM mutation. All samples sequenced and analyzed by Southern analysis have been confirmed as true positives for PSA and PSM.

Experimental Details

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The ability to accurately stage patients with prostate

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cancer at the time of diagnosis is clearly of paramount importance in selecting appropriate therapy and in predicting long-term response to treatment, potential cure. Pre-surgical staging presently consists of physical examination, serum PSA and PAP determinations, and numerous imaging modalities including transrectal ultrasonography, CT scanning, radionuclide bone scans, and even MRI scanning. present modality, however, addresses the issue of hematogenous micrometastatic disease and the potential negative impact on prognosis that this may produce. Previous work has shown that only a percentage of circulating tumor cells will inevitably go on to form a solid metastasis (16), however, the detection of and potential quantification circulating tumor cell burden may prove valuable in more accurately staging disease. The long-term impact of hematogenous micrometastatic disease must be studied by comparing the clinical courses of patients found to have these cells in their circulation with patients of similar stage and treatment who test negatively.

The significantly higher level of detection of tumor cells with PSM as compared to PSA is not surprising to us, since more consistent expression of PSM in prostate carcinomas of all stages and grades as compared to variable expression of PSA in more poorly differentiated and anaplastic prostate cancers is The detection of tumor cells in the three patients that had undergone radical prostatectomies with subsequent undetectable amounts of serum PSA was suprising. These patients would be considered to be surgical "cures" by standard criteria, apparently continue to harbor prostatic tumor cells. It will be interesting to follow the clinical course of these patients as compared to others without PCR evidence of residual disease.

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#### EXAMPLE 4:

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EXPRESSION OF THE PROSTATE SPECIFIC MEMBRANE ANTIGEN

(PSM) DIMINISHES THE MITOGENIC STIMULATION OF

AGGRESSIVE HUMAN PROSTATIC CARCINOMA CELLS BY

TRANSFERRIN

An association between transferrin and human prostate cancer has been suggested by several investigators. It has been shown that the expressed prostatic secretions of patients with prostate cancer are enriched with respect to their content of transferrin and that prostate cancer cells are rich in transferrin receptors (J. Urol. 143, 381, 1990). Transferrin derived from bone marrow has been shown to selectively stimulate the growth of aggressive prostate cancer cells (PNAS 89, 6197, 1992). DNA sequence analysis has revealed that a portion of the coding region, from nucleotide 1250 to 1700 possesses a 54% homology to the human transferrin PC-3 cells do not express PSM mRNA or protein and exhibit increased cell growth in response to transferrin, whereas, LNCaP prostate cancer cells which highly express PSM have a very weak response to To determine whether PSM expression by prostatic cancer cells impacts upon their mitogenic response to transferrin the full-length PSM cDNA was transfected into the PC-3 prostate cancer cells. Clones highly expressing PSM mRNA were identified by Northern analysis and expression of PSM protein was verified by Western analysis using the monoclonal antibody 7E11-C5.3.

 $2 \times 10^4$  PC-3 or PSM-transfected PC-3 cells per well ere plated in RPMI medium supplemented with 10% fetal bovine serum and at 24 hrs. added 1  $\mu$ g per ml. of holotransferrin to the cells. Cells were counted at 1 day to be highly mitogenic to the PC-3 cells. Cells

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were counted at 1 day to determine plating efficiency and at 5 days to determine the effect of the transferrin. Experiments were repeated to verify the results.

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PC-3 cells experienced an average increase of 275% over controls, whereas the LNCaP cells were only stimulated 43%. Growth kinetics revealed that the PSM-transfected PC-3 cells grew 30% slower than native PC-3 cells. This data suggests that PSM expression in aggressive, metastatic human prostate cancer cells significantly abrogates their mitogenic response to transferrin.

The use of therapeutic vaccines consisting of cytokinesecreting tumor cell preparations for the treatment of 15 established prostate cancer was investigated in the Dunning R3327-MatLyLu rat prostatic adenocarcinoma. Only IL-2 secreting, irradiated tumor cell model. preparations were capable of curing animals from subcutaneously established tumors, and engendered 20 immunological memory that protected the animals from Immunotherapy was less another tumor challenge. effective when tumors were induced orthotopically, but nevertheless led to improved outcome, significantly delaying, and occasionally preventing recurrence of 25 tumors after resection of the cancerous prostate. Induction of a potent immune response in tumor bearing animals against the nonimmunogenic MatLyLu tumor supports the view that active immunotherapy of prostate cancer may have therapeutic benefits. 30

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#### EXAMPLE 5:

# CLONING AND CHARACTERIZATION OF THE PROSTATE SPECIFIC MEMBRANE ANTIGEN (PSM) PROMOTER.

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The expression and regulation of the PSM gene is complex. By immunostaining, PSM antigen was found to be expressed brilliantly in metastasized tumor, and in organ confined tumor, less so in normal prostatic tissue and more heterogenous in BPH. PSM is strongly expressed in both anaplastic and hormone refractory tumors. PSM mRNA has been shown to be down regulated Expression of PSM RNA is also modulated by androgen. by a host of cytokines and growth factors. Knowledge of the regulation of PSM expression should aid in such diagnostic and therapeutic strategies imunoscintigraphic imaging of prostate cancer protate-specific promoter-driven gene therapy.

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- Sequencing of a 3 kb genomic DNA clone that contained 2.5 kb upstream of the transcription start site revealed that two stretches of about 300 b.p. (-260 to -600; and -1325 to -1625) have substantial homology (79-87%) to known genes. The promoter lacks a GC rich region, nor does it have a consensus TATA box. However, it contains a TA-rich region from position -35 to -65.
- Several consensus recognition sites for general transcription factors such as AP1, AP2, NFkB, GRE and E2-RE were identified. Chimeric constructs containing fragments of the upstream region of the PSM gene fused to a promoterless chloramphenicol acetyl transferase gene were transfected into, and transiently expressed in LNCaP, PC-3, and SW620 (a colonic cell line). With an additional SV40 enhancer, sequence from -565 to +76

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exhibited promoter activity in LNCaP but not in PC-3 nor in SW620.

#### Materials and Methods

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Cell Lines. LNCaP and PC-3 prostatic carcinoma cell lines (American Type Culture Collection) were cultured in RPMI and MEM respectively, supplemented with 5% fetal calf serum at 37°C and 5% CO<sub>2</sub>. SW620, a colonic cell line, is a gift from Melisa.

Polymerase Chain Reaction. The reaction was performed in a 50 μl volume with a final concentration of the following reagents: 16.6 mM NH<sub>4</sub>SO<sub>4</sub>, 67 mM Tris-HCl pH 8.8, acetylated BSA 0.2 mg/ml, 2mM MgCl<sub>2</sub>, 250μM dNTPs, 10 mM β-mercaptoethanol, and 1 U of rth 111 Taq polymerase (Boehringer Mannhiem, CA). A total of 25 cycles were completed with the following profile: cycle 1, 94°C 4 min.; cycle 2 through 25, 94°C 1 min, 60°C 1 min, 72°C 1 min. The final reaction was extended for 10 min at 72°C. Aliquots of the reaction were electrophoresed on 1 % agarose gels in 1X Tris-acetate-EDTA buffer.

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Cloning of PSM promoter. A bacteriophage P1 library of human fibroblast genomic DNA (Genomic Sysytems, Inc., St. Louis, MI), was screened using a PCR method of Primers located at the 5' end of PSM Pierce et al. used:5'-CTCAAAAGGGGCCGGATTTCC-3' cDNA were 5'CTCTCAATCTCACTAATGCCTC-3'. A positive clone, was digested with Xhol restriction enzyme. Southern analysis of the restricted fragments using a DNA probe from the extreme 5' to the Ava-1 site of PSM cDNA fragment contains confirmed that a 3Kb regulatory sequence of the PSM gene. The 3 kb Xho1 fragment was subcloned into pKSBluescrpt vectors and

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sequenced using the dideoxy method.

Functional Assay of PSM Promoter. Chloramphenicol Transferase, Acetyl (CAT) gene plasmids were constructed from the Smal-HindIII fragments or subfragements (using either restriction subfragments or PCR) by insertion into promoterless pCAT basic or pCAT-enhancer vectors (Promega). constructs were cotransfected with pSVBgal plasmid (5 μg of each plasmid) into cell lines in duplicates, calcium а phosphate method (Gibco-BRL. Gaithersburg, MD). The transfected cells harvested 72 hours later and assayed (15 $\mu$ g of lysate) for CAT activity using the LSC method and for Bgal activity (Promega). CAT activities were standardized by comparision to that of the Egal activities.

## Results

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20 Sequence of the 5' end of the PSM gene.

The DNA sequence of the 3 kb XhoI fragment of p683 which includes 500 bp of DNA from the RNA start site was determined (Figures 31A-31D) Sequence 683XFRVS starts from the 5' distal end of PSM promoter, it overlaps with the published PSM putative promoter at nt 2485, i.e. the putative transcription start site is at nt 2485; sequence 683XF107 is the reverse, complement of 683XFRVS). The sequence from the XhoI fragment displayed a remarkable arrays of elements and motifs which are characteristic of eukaryotic promoters and regulatory regions found in other genes (Figure 32).

Functional Analysis of upstream PSM genomic elements for promoter activity.

Various pCAT-PSM promoter constructs were tested for promoter activities in two prostatic cell lines:

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LNCaP, PC-3 and a colonic SW620 (Figure 33). Induction of CAT activity was neither observed in p1070-CAT which contained a 1070 bp PSM 5' promoter fragment, nor in p676-CAT which contained a 641 bp PSM 5' promoter fragment. However, with an additional SV-40 enhancer, sequence from -565 to +76 (p676-CATE) exhibited promoter activity in LNCaP but not in PC-3 nor in SW620.

Therefore, a LNCaP specific promoter fragment from -565 to +76 has been isolated which can be used in PSM promoter-driven gene therapy.

#### EXAMPLE 6:

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ALTERNATIVELY SPLICED VARIANTS OF PROSTATE SPECIFIC MEMBRANE ANTIGEN RNA: RATIO OF EXPRESSION AS A

POTENTIAL MEASUREMENT OF PROGRESSION

## 20 MATERIALS AND METHODS

Cell Lines. LNCaP and PC-3 prostatic carcinoma cell lines were cultured in RPMI and MEM respectively, supplemented with 5% fetal calf serum at 37°C and 5% CO<sub>2</sub>.

Primary tissues. Primary prostatic tissues were obtained from MSKCC's in-house tumor procurement service. Gross specimen were pathologically staged by MSKCC's pathology service.

isolated by a Total RNA was RNA Isolation. thiocynate/phenol/chloroform guanidinium modified method using a RNAzol B kit (Tel-Test, Friendswood, RNA was stored in diethyl pyrocarbonate-treated TX). quantified using was at -80°C. RNA water spectrophometric absorption at 260nm.

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cDNA synthesis. Two different batches of normal prostate mRNAs obtained from trauma-dead males (Clontech, Palo Alto, CA) were denatured at 70°C for 10 min., then reverse transcribed into cDNA using random hexamers and Superscript II reverse transcriptase (GIBCO-BRL, Gaithersburg, MD) at 50°C for 30 min. followed by a 94°C incubation for 5 min.

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Polymerase Chain Reaction. Oligonucleotide primers(5'-CTCAAAAGGGGCCGGATTTCC-3' 10 AGGCTACTTCACTCAAAG-3'), specific for the 5' and 3' ends of PSM cDNA were designed to span the cDNA sequence. The reaction was performed in a 50  $\mu$ l volume with a final concentration of the following reagents: 16.6 mM NH,SO,, 67 mM Tris-HCl pH 8.8, acetylated BSA 0.2 mg/ml, 15 2mM MgCl<sub>2</sub>, 250 $\mu$ M dNTPs, 10 mM ß-mercaptoethanol, and 1 U of rTth polymerase (Perkin Elmer, Norwalk, CT). total of 25 cycles were completed with the following profile: cycle 1, 94°C 4 min.; cycle 2 through 25, 94°C 20 1 min, 60°C 1 min, 72°C 1 min. The final reaction was extended for 10 min at 72°C. Aliquots of the reaction were electrophoresed on 1 % agarose gels in 1X Trisacetate-EDTA buffer.

- 25 Cloning of PCR products. PCR products were cloned by the TA cloning method into pCRII vector using a kit from Invitrogen (San Diego, CA). Ligation mixture were transformed into competent Escherichia coli Inv5α.
- Sequencing. Sequencing was done by the dideoxy method using a sequenase kit from US Biochemical (Cleveland, OH). Sequencing products were electrophoresed on a 5% polyacrylamide/7M urea gel at 52°C.
  - RNase Protection Assays. Full length PSM cDNA clone was digested with NgoM 1 and Nhel. A 350 b.p. fragment

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was isolated and subcloned into pSPORT1 vector (GIBCO-BRL, Gaithersburg, MD). The resultant plasmid, pSP350, was linearized, and the insert was transcribed by SP6 RNA polymerase to yield antisense probe of 395 nucleotide long, of which 355 nucleotides and/or 210 nucleotides should be protected from RNAse digestion by PSM or PSM' RNA respectively (Fig.2). Total celluar RNA (20  $\mu$ g) from different tissues were hybridized to the aforementioned antisense RNA probe. Assays were performed as described (7). tRNA was used as negative control. RPAs for LNCaP and PC-3 were repeated.

#### RESULTS

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RT-PCR of mRNA from normal prostatic tissue. Two independent RT-PCR of mRNA from normal prostates were 15 performed as described in Materials and Methods. Subsequent cloning and sequencing of the PCR products revealed the presence of an alternatively spliced PSM' has a shorter cDNA (2387 PSM'. variant, nucleotides) than PSM (2653 nucleotides). The results 20 of the sequence analysis are shown in Figure 34. cDNAs are identical except for a 266 nucleotide region near the 5' end of PSM cDNA (nucleotide 114 to 380) is absent in PSM' cDNA. Two independent repetitions of RT-PCR of different mRNA samples yielded 25 identical results.

RNase Protection Assays. An RNA probe complementary to PSM RNA and spanning the 3' splice junction of PSM' RNA was used to measure relative expression of PSM and PSM' mRNAs (Figure 35). With this probe, both PSM and PSM' RNAs in LNCaP cells was detected and the predominant form was PSM. Neither PSM nor PSM' RNA was detected in PC-3 cells, in agreement with previous Northern and Western blot data (5,6). Figure 36 showed the presence of both splice variants in human primary prostatic tissues. In primary prostatic tumor, PSM is

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the dominant form. In contrast, normal prostate expressed more PSM' than PSM. BPH samples showed about equal expression of both variants.

Tumor Index. The relative expression of PSM and PSM' (Figure 36) was quantified by densitometry and expressed as a tumor index (Figure 37). LNCaP has an index ranging from 9-11; CaP from 3-6; BPH from 0.75 to 1.6; normal prostate has values from 0.075 to 0.45.

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#### DISCUSSION

Sequencing data of PCR products derived from human normal prostatic mRNA with 5' and 3' end PSM oligonucleotide primers revealed a second splice variant, PSM', in addition to the previously described PSM cDNA.

PSM is a 750 a.a. protein with a calculated molecular weight of 84,330. PSM was hypothesized to be a type II integral membrane protein (5). A classic type II membrane protein is the transferrin receptor and indeed PSM has a region that has modest homology with the transferrin receptor (5). Analysis of the PSM amino acid sequence by either the methods of Rao and Argos (7) or Eisenburg et. al. (8) strongly predicted one transmembrane helix in the region from a.a.#20 to #43. Both programs found other regions that could be membrane associated but were not considered likely candidates for being transmembrane regions.

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PSM' antigen, on the other hand, is a 693 a.a. protein as deduced from its mRNA sequence with a molecular weight of 78,000. PSM' antigen lacks the first 57 amino acids present in PSM antigen (Figure 34). It is likely that PSM' antigen is cytosolic.

The function of PSM and PSM' are probably different.

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The cellular location of PSM antigen suggests that it may interact with either extra- or intra- cellular ligand(s) or both; while that of PSM' implies that PSM' can only react with cytosolic ligand(s). Furthermore, PSM antigen has 3 potential phosphorylation sites on These sites are absent in PSM' its cytosolic domain. On the other hand, PSM' antigen has 25 antigen. potential phosphorylation sites, 10 N-myristoylation sites and 9 N-glycosylation sites. For PSM antigen, all of these potential sites would be on The modifications of these extracellular surface. sites for these homologous proteins would be different depending on their cellular locations. Consequently, the function(s) of each form would depend on how they are modified.

The relative differences in expression of PSM and PSM'. by RNase protection assays was analyzed. Results of expression of PSM and PSM' in primary prostatic tissues strongly suggested a relationship between the relative expression of these variants and the status of the While it is noted cell: either normal or cancerous. here that the sample size of the study is small (Figures 36 and 37), the consistency of the trend is The samples used were gross specimens from evident. The results may have been even more dramatic if specimens that were pure in content of CaP, BPH or Nevertheless, in these normal had been used. specimens, it is clear that there is a relative increase of PSM over PSM' mRNA in the change from 30 normal to CaP. The Tumor Index (Figure 37) could be useful in measuring the pathologic state of a given It is also possible that the change in sample. expression of PSM over PSM' may be a reason for tumor progression. A more differentiated tumor state may be 35 restored by PSM' either by transfection or by the use of differentiation agents.

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#### EXAMPLE 7:

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ENHANCED DETECTION OF PROSTATIC HEMATOGENOUS MICRO-METASTASES WITH PSM PRIMERS AS COMPARED TO PSA PRIMERS USING A SENSITIVE NESTED REVERSE TRANSCRIPTASE-PCR ASSAY.

randomly selected samples were analyzed from 77 patients with prostate cancer and reveals that PSM and PSA primers detected circulating prostate cells in 48 (62.3%) and 7 (9.1%) patients, respectively. treated stage D disease patients, PSM primers detected cells in 16 of 24 (66.7%), while PSA primers detected cells in 6 of 24 patients (25%). In hormone-refractory prostate cancer (stage D3), 6 of 7 patients were positive with both PSA and PSM primers. All six of these patients died within 2-6 months of their assay, despite aggressive cytotoxic chemotherapy, in contrast to the single patient that tested negatively in this group and is alive 15 months after his assay, suggesting that PSA-PCR positivity may serve as a predictor of early mortality. In post-radical prostatectomy patients with negative serum PSA values, psm primers detected metastases in 21 of 31 patients (67.7%), while PSA primers detected cells in only 1 of 33 (3.0%), indicating that micrometastatic spread may be a relatively early event in prostate cancer. analysis of 40 individuals without known prostate cancer provides evidence that this assay is highly specific and suggests that PSM expression may predict in patients without development of cancer clinically apparent prostate cancer. Using PSM primers, micrometastases were detected in 4 of 40 controls, two of whom had known BPH by prostate biopsy and were later found to have previously undetected prostate cancer following repeat prostate biopsy

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performed for a rising serum PSA value. These results show the clinical significance of detection of hematogenous micrometastatic prostate cells using PSM primers and potential applications of this molecular assay.

#### EXAMPLE 8:

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# MODULATION OF PROSTATE SPECIFIC MEMBRANE ANTIGEN (PSM) EXPRESSION IN VITRO BY CYTOKINES AND GROWTH FACTORS.

The effectiveness of CYT-356 imaging is enhanced by manipulating expression of PSM. PSM mRNA expression is downregulated by steroids. This is consistent with the clinical observations that PSM is strongly expressed in both anaplastic and hormone refractory lesions. In contrast, PSA expression is decreased following hormone. withdrawal. In hormone refractory disease, believed that tumor cells may produce both growth factors and receptors, thus establishing an autocrine loop that permits the cells to overcome normal growth Many prostate tumor epithelial cells constraints. express both  $TGF\alpha$  and its receptor, epidermal growth. factor receptor. Results indicate that the effects of  $TGF\alpha$  and other selected growth factors and cytokines on the expression of PSM in-vitro, in the human prostatic carcinoma cell line LNCaP.

2x10<sup>6</sup> LNCaP cells growing in androgen-depleted media were treated for 24 to 72 hours with EGF, TGFα, TNFß or TNFα in concentrations ranging from 0.1 ng/ml to 100 ng/ml. Total RNA was extracted from the cells and PSM mRNA expression was quantitated by Northern blot analysis and laser densitometry. Both b-FGF and TGFα yielded a dose-dependent 10-fold upregulation of PSM expression, and EGF a 5-fold upregulation, compared to untreated LNCaP. In contrast, other groups have shown

a marked downregulation in PSA expression induced by these growth factors in this same in-vitro model.  $TNF\alpha$ , which is cytotoxic to LNCaP cells, and  $TNF\beta$  downregulated PSM expression 8-fold in androgen depleted LNCaP cells.

TGF $\alpha$  is mitogenic for aggressive prostate cancer cells. There are multiple forms of PSM and only the membrane form is found in association with tumor progression. The ability to manipulate PSM expression by treatment with cytokines and growth factors may enhance the efficacy of Cytogen 356 imaging, and therapeutic targeting of prostatic metastases.

#### 15 **EXAMPLE 9:**

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NEOADJUVANT ANDROGEN-DEPRIVATION THERAPY (ADT) PRIOR TO RADICAL PROSTATECTOMY RESULTS IN A SIGNIFICANTLY DECREASED INCIDENCE OF RESIDUAL MICROMETASTATIC DISEASE AS DETECTED BY NESTED RT-PCT WITH PRIMERS.

Radical prostatectomy for clinically localized prostate cancer is considered by many the "gold standard" treatment. Advances over the past decade have served to decrease morbidity dramatically. Improvements intended to assist clinicians in better staging patients preoperatively have been developed, however the incidence of extra-prostatic spread still exceeds 50%, as reported in numerous studies. A phase III prospective randomized clinical study designed to compare the effects of ADT for 3 months in patients undergoing radical prostatectomy with similarly matched controls receiving surgery alone was conducted. previously completed phase II study revealed a 10% margin positive rate in the ADT group (N=69) as compared to a 33% positive rate (N=72) in the surgery alone group.

Patients who have completed the phase III study were analyzed to determine if there are any differences between the two groups with respect to residual micrometastatic disease. A positive PCR result in a post-prostatectomy patient identifies viable metastatic cells in the circulation.

Nested RT-PCR was performed with PSM primers on 12 patients from the ADT group and on 10 patients from the control group. Micrometastatic cells were detected in 10 9/10 patients (90%) in the control group, as compared to only 2/12 (16.7%) in the ADT group. In the ADT group, 1 of 7 patients with organ-confined disease tested positively, as compared to 3 of 3 patients in the control group. In patients with extra-prostatic 15 disease, 1 of 5 were positive in the ADT group, as compared to 6 of 7 in the control group. These results indicate that a significantly higher number of patients may be rendered tumor-free, and potentially "cured" by the use of neoadjuvant ADT. 20

## EXAMPLE 10:

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# SENSITIVE NESTED RT-PCR DETECTION OF CIRCULATION PROSTATIC TUMOR CELLS - COMPARISON OF PSM AND PSA-BASED ASSAYS

Despite the improved and expanded arsenal of modalities available to clinician today, including sensitive serum PSA assays, CT scan, transrectal ultrasonography, endorectal co.I MRI, etc., many patients are still found to have metastatic disease at the time of pelvic lymph node dissection and radical prostatectomy. A highly sensitive reverse transcription PCR assay capable of detecting occult hematogenous micrometastatic prostatic cells that would otherwise go undetected by presently available staging modalities

was developed. This assay is a modification of similar PCR assays performed in patients with prostate cancer and other malignancies<sup>2,3,4,5</sup>. The assay employs PCR primers derived from the cDNA sequences of prostate-specific antigen<sup>6</sup> and the prostate-specific membrane antigen recently cloned and sequenced.

## Materials and Methods

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LNCaP and MCF-7 cells were Cells and Reagents. 10 obtained from the American Type Culture Collection (Rockville, MD.). Details regarding the establishment and characteristics of these cell lines have been previously published<sup>8,9</sup>. Cells grown in RPMI 1640 medium and supplemented with L-glutamine, nonessential 15 amino acids, and 5% fetal calf serum (Gibco-BRL, In a 5% CO, incubator at 37°C. Gaithersburg, MD.) media was obtained from the MSKCC Media Preparation Facility. Routine chemical reagents were of the highest grade possible and were obtained from 20 Sigma Chemical Company (St. Louis, MO).

> Patient Blood Specimens. All blood specimens used in this study were from patients seen in the outpatient offices of urologists on staff at MSKCC. Two anticoagulated tubes per patient were obtained at the time of their regularly scheduled blood draws. were obtained with informed consent of each patient , as per a protocol approved by the MSKCC Institutional Samples were promptly brought to the Review Board. laboratory for immediate processing. Seventy-seven specimens from patients with prostate cancer were randomly selected and delivered to the laboratory "blinded" along with samples from negative controls for These included 24 patients with stage D processing. disease (3 with  $D_0$ , 3 with  $D^1$ , 11 with  $\overrightarrow{D}$ , and 7 with  ${ ilde D}^3)$ , 31 patients who had previously undergone radical

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prostatectomy and had undetectable postoperative serum PSA levels (18 with pT2 lesions, 11 with pT3, and 2 pT4), 2 patients with locally recurrent disease following radical prostatectomy, 4 patients who had received either external beam radiation therapy or interstitial l125 implants, 10 patients with untreated clinical stage T1-T2 disease, and 6 patients with clinical stage T3 disease on anti-androgen therapy. The forty blood specimens used as negative controls were from 10 health males, 9 males with biopsy-proven 10 BPH and elevated serum PSA levels, 7 healthy females, 4 male patients with renal cell carcinoma, 2 patients with prostatic intraepithelial neoplasia (PIN), patients with transitional cell carcinoma of the bladder and a pathologically normal prostate, 1 patient 15 patient with acute prostatitis, 1 promyelocytic leukemia, 1 patient with testicular cancer, 1 female patient with renal cell carcinoma, 1 patient with lung cancer, and 1 patient with a cyst of the testicle. 20

> Blood Sample Processing/RNA Extraction. 4 ml of whole anticoagulated venous blood was mixed with 3 ml of ice cold PBS and then carefully layered atop 8 ml of Ficoll (Pharmacia, Uppsala, Sweden) in a 14-ml polystyrene Tubes were centrifuged at 200 x g for 30 min. at tube. The buffy coat layer (approx. 1 ml.) was 4°C. carefully removed and rediluted to 50 ml with ice cold PBS in a 50 ml polypropylene tube. This tube was then centrifuged at 2000 x g for 30 min. at 4°C. supernatant was carefully decanted and the pellet was allowed to drip dry. One ml of RNazol B was then added to the pellet and total RNA was isolated as per manufacturers directions (Cinna/Biotecx, Houston, TX.) RNA concentrations and purity were determined by UV spectroscopy on a Beckman DU 640 spectrophotometer and by gel analysis.

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Determination of PCR Sensitivity. RNA was isolated from LNCaP cells and from mixtures of LNCaP and MCF-7 cells at fixed ratios (i.e. 1:100, 1:1,000, etc.) using RNAzol B. Nested PCR was then performed as described below with both PSA and PSM primers in order to determine the limit of detection for the assay. LNCaP:MCF-7 (1:100,000) cDNA was diluted with distilled water to obtain concentrations of 1:1,000,000. The human breast cancer cell line MCF-7 was chosen because they had previously been tested by us and shown not to express either PSM nor PSA by both immunohistochemistry and conventional and nested PCR.

Polymerase Chain Reaction. The PSA outer primer sequences are nucleotides 494-513 (sense) in exon 4 and nucleotides 960-979 (anti-sense) in exon 5 of the PSA cDNA. These primers yield a 486 bp PCR product from PSA CDNA that can be distinguished from a product synthesized from possible contaminating genomic DNA.

PSA-494 5'-TAC CCA CTG CAT CAG GAA CA-3' PSA-960 5'-CCT TGA AGC ACA CCA TTA CA-3'

The PSA inner upstream primer begins at nucleotide 559 and the downstream primer at nucleotide 894 to yield a 355 bp PCR product.

PSA-559 5'-ACA CAG GCC AGG TAT TTC AG-3' PSA-894 5'-GTC CAG CGT CCA GCA CAC AG-3'

the synthesized by All primers were Microchemistry Core Facility. 5μg of total RNA was reverse-transcribed into cDNA using random hexamer and Superscript II reverse (Gibco-BRL) to the (Gibco-BRL) according transcriptase manufacturers recommendations.  $1\mu l$  of this CDNA served as the starting template for the outer primer PCR The  $20\mu$ l PCR mix included: 0.5U Tag reaction. polymerase (Promega) Promega reaction buffer, 1.5mM  $MgCl_2$ , 200 $\mu$ M dNTPs, and 1.0 $\mu$ M of each primer. This mix was then transferred to a Perkin Elmer 9600 DNA thermal cycler and incubated for 25 cycles. The PCR profile was as follows: 94°C x 15 sec., 60°C x 15 sec., and 72°C for 45 sec. After 25 cycles, samples were placed on ice, and  $1\mu$ l of this reaction mix served as the template for another 25 cycles using the inner primers. The first set of tubes were returned to the thermal cycler for 25 additional cycles. The PSM outer upstream primer sequences are nucleotides 1368-1390 and the downstream primers are nucleotides 1995-2015, yielding a 67 bp PCR product.

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PSM-1368 5'-CAG ATA TGT CAT TCT GGG AGG TC-3' PSM-2015 5'-AAC ACC ATC CCT CGA ACC-3'

The PSM inner upstream primer span nucleotides 1689-1713 and the downstream primer span nucleotides 1899-1923, yielding a 234 bp PCR product.

PSM-1689 5'-CCT AAC AAA AGA GCT GAA AAG CCC-3' PSM-1923 5'-ACT GTG ATA CAG TGG ATA GCC GCT-3'

 $2\mu l$  of cDNA was used as the starting DNA template in 20 The  $50\mu l$  PCR mix included: 1U Taq the PCR assay. polymerase (Boehringer Mannheim), 250 µM cNTPs, 10 mM ßmercaptoethanol, 2mM MgCl, and  $5\mu$ l of a 10x buffer mix containing: 166mM NH<sub>2</sub>SO<sub>4</sub>, 670mM Tris pH 8.8, and 2mg/ml PCR was carried out in a Perkin of acetylated BSA. 25 Elmer 480 DNA thermal cycler with the following parameters: 94°C x 4 minutes for 1 cycle, 94°C x 30 sec., 58°C x 1 minute, and 72°C x 1 minute for 25 Samples were cycles, followed by 72°C x 10 minutes. then iced and  $2.5\mu l$  of this reaction mix was used as 30 the template for another 25 cycles with a new reaction mix containing the inner PSM primers. cDNA quality was verified by performing control reactions using primers derived from the ß-2-microglobulin gene sequence 10 a ubiquitous housekeeping gene. These primers span exons 35 2-4 and generate a 620 bp PCR product. The sequences for these primers are:

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B-2 (exon 2) 5'-AGC AGA GAA TGG AAA GTC AAA-3' B-2 (exon 4) 5'-TGT TGA TGT TGG ATA AGA GAA-3' The entire PSA mix and 7-10µl of each PSM reaction mix were run on 1.5-2% agarose gels, stained with ethidium bromide and photographed in an Eage Eye Video Imaging System (Statagene, Torrey Pines, CA.). Assays were repeated at least twice to verify results.

Cloning and Sequencing of PCR Products. PCR products were cloned into the pCR II plasmid vector using the TA 10 cloning system (Invitrogen). These plasmids were transformed into competent E. coli cells using standard methods 11 and plasmid DNA was isolated using Magic and screened by restriction Minipreps (Promega) Double-stranded TA clones 15 sequenced by the dideoxy method 12 using 35S-cCTP (NEN) and Sequenase (U.S. Biochemical). Sequencing products were then analyzed on 6% polyacrilamide/7M urea gels, which were fixed, dried, and autoradiographed as 20 described.

> Southern Analysis. PCR products were transferred from ethidium-stained agarose gels to Nytran nylon membranes (Schletcher and Schuell) by pressure blotting with a (Stratagene) according Posi-blotter DNA was cross-linked to manufacturer's instructions. the membrane using a UV Stratalinker (Stratagene). Blots were pre-hybridized at 65°C for 2 hours and subsequently hybridized with denatured 32P-labeled, random-primed 13 cDNA probes (either PSA or PSM). Blots were washed twice in 1x SSC/0.5% SDS at 42°C and twice in 0.1x SSC/0.1% SDS at 50°C for 20 minutes each. Membranes were air-dried and autoradiographed for 1-3 hours at room temperature with Hyperfilm MP (Amersham).

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#### Results

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PSA and PSM Nested PCR Assays: The application of nested PCR increased the level of detection from an average of 1:10,000 using outer primers alone, better than 1:1,000,000. Dilution curves demonstrating this added sensitivity are shown for PSA and PSM-PCR in Figures 1 and 2 respectively. Figure 1 shows that the 486 bp product of the PSA outer primer set is clearly to staining with ethidium detectable dilutions, whereas the PSA inner primer 355 bp product is clearly detectable in all dilutions shown. Figure 2 the PSM outer primer 647 bp product is also clearly detectable in dilutions to only 1:10,000 with conventional PCR, in contrast to the PSM inner nested PCR 234 bp product which is detected in dilutions as low as 1:1,000,000. Southern blotting was performed on all controls and most of the patient samples in order Southern blots of confirm specificity. respective dilution curves confirmed the primer specificities but did not reveal any significantly increased sensitivity.

performed on 40 samples from patients and volunteers as described in the methods and materials section. Figure 48 reveals results from 4 representative negative control specimens, in addition to a positive control. Each specimen in the study was also assayed with the ß-2-microglobulin control, as shown in the figure, in order to verify RNA integrity. Negative results were obtained on 39 of these samples using the PSA primers, however PSM nested PCR yielded 4 positive results. Two of these "false positives" represented patients with elevated serum PSA values and an enlarged prostate who underwent a transrectal prostate biopsy revealing stromal and fibromuscular hyperplasia. In both of

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these patients the serum PSA level continued to rise and a repeat prostate biopsy performed at a later date revealed prostate cancer. One patient who presented to the clinic with a testicular cyst was noted to have a positive PSM nested PCR result which has been unable to Unfortunately, this patient never returned for follow up, and thus have not been able to obtain another blood sample to repeat this assay. Positive result were obtained with both PSA and PSM primers in a 61 year old male patient with renal cell carcinoma. This patient has a normal serum PSA level and a normal digital rectal examination. Overall, if the two patients were excluded in whom a positive PCR, but no other clinical test, accurately predicted the presence of prostate cancer, 36/38 (94.7%) of the negative controls were negative with PSM primers, and 39/40 (97.5%) were negative using PSA primers.

Patient Samples: In a "blinded" fashion, in which the laboratory staff were unaware of the nature of each specimen, 117 samples from 77 patients mixed randomly with 40 negative controls were assayed. The patient samples represented a diverse and heterogeneous group as described earlier. Several representative patient samples are displayed in Figure 49, corresponding to positive results from patients with both localized and disseminated disease. Patients 4 and 5, both with stage D prostate cancer exhibit positive results with both the outer and inner primer pairs, indicating a large circulating tumor cell burden, as compared to the Although the PSM and PSA primers other samples. yielded similar sensitivities in LNCaP dilution curves primers previously shown. PSM as micrometastases in 62.3% of the patient samples, whereas PSA primers only detected 9.1%. In patients with documented metastatic prostate cancer (stages Dn receiving anti-androgen treatment, PSM primers

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detected micrometastases in 16/24 (66.7%), whereas PSA primers detected circulating cells in only 6/24 (25%). In the study 6/7 patients with hormone-refractory prostate cancer (stage D<sub>3</sub>) were positive. In the study, PSA primers revealed micrometastatic cells in only 1/15 (6.7%) patients with either pT3 or pT4 (locally-advanced) prostate cancer following radical prostatectomy. PSM primers detected circulating cells in 9/15 (60%) of these patients. Interestingly, circulating cells 13/18 (72.2%) patients with pT2 (organ-confined) prostate cancer following radical prostatectomy using PSM primers was detected. None of these patient samples were positive by PSA-PCR.

Improved and more sensitive method for the detection of minimal, occult micrometastic disease have been reported for a number of malignancies by use of immunohistochemical methods (14), as well as the polymerase chain reaction (3, 4, 5). The application of PCR to detect occult hematogenous micrometastases in prostate cancer was first described by Moreno, et al. (2) using conventional PCR with PSA-derived primers.

When human prostate tumors and prostate cancer cells in-vitro were studied by immunohistochemistry and mRNA analysis, PSM appeared to be highly expressed in anaplastic cells, hormone-refractory cells, and bony metastases (22, 23, 24), in contrast to PSA. If cells capable of hematogenous micrometastasis represent the more aggressive and poorly-differentiated cells, they may express a higher level of PSM per cell as compared to PSA, enhancing their detectibility by RT-PCR.

Nested RT-PCR assays are both sensitive and specific.

Results have been reliably reproduced on repeated occasions. Long term testing of both cDNA and RNA stability is presently underway. Both assays are

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capable of detecting one prostatic cell in at least one million non-prostatic cells of similar size. confirms the validity of the comparison of PSM vs. PSA Similar levels of PSM expression in both human prostatic cancer cells in-vivo and LNCaP cells in-vitro resulted. The specificity of the PSM-PCR assay was supported by the finding that two "negative control" patients with positive PSM-PCR results were both subsequently found to have prostate cancer. suggests an exciting potential application for this technique for use in cancer screening. In contrast to recently published data (18), significant ability for PSA primers to accurately detect micrometastatic cells in patients with pathologically with pathologically organ-confined prostate cancer, despite the sensitivity of the assay failed to result. Rather a surprisingly high percentage of patients with localized prostate cancer that harbor occult circulating prostate cells following "curative" radical prostatectomy results which suggests that micrometastasis is an early event in prostate cancer.

The application of this powerful new modality to potentially stage and/or follow the response to therapy in patients with prostate cancer certainly merits further investigation. In comparison to molecular detection of occult tumor cells, present clinical modalities for the detection of prostate cancer spread appear inadequate.

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#### EXAMPLE 11:

# CHROMOSOMAL LOCALIZATION OF COSMID CLONES 194 AND 683 BY FLUORESCENCE IN-SITU HYBRIDIZATION:

PSM was initially mapped as being located on chromosome 11pll.2-pl3 (Figures 51-54). Further information from CDNA in-situ hybridizations the experiments demonstrated as much hybridization on the q as p arms. Much larger fragments of genomic DNA was obtained as cosmids and two of these of about 60 kilobases each one going 3' and the other 5' both demonstrated binding to chromosome 11 p and q under low stringency. under higher stringency conditions only the binding at 11q14-q21 remained. This result suggests that there is another gene on 11p that is very similar to PSM because it is so strongly binding to nearly 120 kilobases of genomic DNA (Figure 50).

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Purified DNA from cosmid clones 194 and 683 was labelled with biotin dUTP by nick translation. Labelled probes were combined with sheared human DNA and independently hybridized to normal metaphase chromosomes derived from PHA stimulated peripheral lymphocytes in a solution containing 50% formamide, 10% dectran sulfate, and 2XSSC. hybridization signals were detected by incubating the hybridized slides in fluoresein conjugated avidin. signal detection the slides Following counterstained with propidium iodide and analyzed. These first experiments resulted in the specific labelling of a group C chromosome on both the long and This chromosome was believed to be chromosome 11 on the basis of its size and morphology. A second set of experiments were performed in which a centromere specific probe 11 chromosome was cohybridized with the cosmid clones. These experiments were carried out in 60% formamide in an attempt to eliminate the cross reactive signal which was observed when low stringency hybridizations were done. These experiments resulted in the specific labelling of the centromere and the long arm of chromosome 11. Measurements of 10 specifically labelled chromosomes 11 demonstrated that the cosmid clones are located at a position which is 44% of the distance from the centromere to the telomere of chromosome arm 11q, an area that corresponds to band 14q. A total of 160 metaphase cells were examined with 153 cells exhibiting specific labelling.

Cloning of the 5' upstream and 3' downstream regions of the PSM genomic DNA. A bacteriophage P1 library of human fibroblast genomic DNA (Genomic Systems, St. Louis, MI) was screened using the PCR method of Pierce et. al. Primer pairs located at either the 5' or 3' termini of PSM cDNA were used. Positive cosmid clones were digested with restriction enzymes and confirmed by Southern analysis using probes which were constructed from either the 5' or 3' ends of PSM cDNA. Positive clone p683 contains the 5' region of PSM cDNA and about 60 kb upstream region. Clone -194 contains the 3' terminal of the PSM cDNA and about 60 kb downstream.

#### EXAMPLE 12:

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# PEPTIDASE ENZYMATIC ACTIVITY

PSM is a type two membrane protein. Most type two membrane proteins are binding proteins, transport proteins or peptidases. PSM appears to have peptidase activity. When examining LNCaP cells with a substrate N-acetyl-aspartyl-14C-glutamic acid, NAAG, glutamic acid was released, thus acting as a carboxypeptidase. In

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vitro translated PSM message also had this peptidase activity..

The result is that seminal plasma is rich in its content of glutamic acid, and are able to design 5 inhibitors to enhance the activity of the non degraded normal substrate if its increased level will have a biologic desired activity. Also biologic activity can be measured to see how it correlates wit the level of 10 Tissue may be examined for activity directly rather than indirectly using in-situ analysis immunohistochemical probes. Because there is another gene highly similar on the other arm of chromosome 11 when isolated the expressed cloned genes can be used to determine what are the substrate differences and use 15 those substrates for identification of PSM related activity, say in circulating cells when looking for metastases.

#### 20 **EXAMPLE 13:**

# IONOTROPICGLUTAMATE RECEPTOR DISTRIBUTION IN PROSTATE TISSUE

#### 25 Introduction:

Excitatory neurotransmission in the central nervous system (CNS) is mediated predominantly by glutamate receptors. Two types of glutamate receptors have been identified in human CNS: metabotropic receptors, which are coupled to second-messenger systems, and ionotropic receptors, which serve as ligand-gated ion channels. The presence of ionotropic glutamate receptors in human prostate tissue was investigated.

#### 35 Methods:

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Detection of glutamate receptor expression was performed using anti-GluR2/3 and anti-biotin

immunohistochemical technique in paraffin-embedded antigen tissues. PSM prostate human neurocarboxypeptidase that acts to release glutamate. In the CNS glutamate acts as a neurotransmitter by acting on glutaminergic ion channels and increases the flow of ions like calcium ions. One way the glutamate signal is transduced into cell activity is activation of nitric oxide synthase, and nitric oxide synthase has recently been found to be present in human prostatic tissue. NO is a major signalling mechanism and is involved in control of cell growth and death, in response to inflammation, in smooth muscle cell contraction, etc,. In the prostate much of the stroma is smooth muscle. It was discovered that the prostate is rich in glutaminergic receptors and have begun to Stromal abnormalities are define this relationship. Stromal epithelial BPH. key feature of interactions are of importance in bothe BPH and CaP. The other glutaminergic receptors through G proteins to change the metabolism of the cell.

#### Results:

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Anti-GluR2/3 immunoreactivity was unique to prostatic stroma and was absent in the prostatic epithelial compartment. Strong anti-GluR4 immunoreactivity was observed in basal cells of prostatic acini.

#### Discussion:

The differential distribution of ionotropic glutamate receptor subtypes between the stromal and epithelial compartments of the prostate has not been previously described. Prostate-specific membrane antigen (PSMA) has an analogous prostatic distribution, with expression restricted to the epithelial compartment.

PSM antigen is a neurocarboxypeptidase that acts to

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-135-

release glutamate from NAAG 1, also a potential In the CNS glutamate acts as a nerotransmitter. neurotransmitter by acting on glutaminergic channels and increases the flow of ions like calcium One way the glutamate signal is transduced into cell activity is the activation of nitric oxide synthase, and nitric oxide synthase has recently been found to be present in human prostatic tissue. NO is a major signaling mechanism and is involved in control of cell growth and death, in response to inflammation, in smooth muscle cell contraction, etc... In the prostate much of the stroma is smooth muscle. The prostate is rich in glutaminergic receptors. abnormalities are the key feature of BPH. epithelial interactions are of importance in both BPH The other glutaminergic receptors through G proteins to change the metabolism of the cell. Glutamate can be produced in the cerebral cortex through the carboxypeptidase activity of the prostatespecific membrane antigen (PSMA). In this location, PSMA cleaves glutamate from acetyl-aspartyl-glutamate. Taken together, these observations suggest a function for PSMA in the human prostate; glutamate may be an autocrine and/or paracrine signalling possibly mediating epithelial-stromal interactions. Ionotropic glutamate receptors display a unique compartmental distribution in the human prostate.

The carboxypeptidase like activity and one substrate is the dipeptide N-acetyl-aspartyl glutamic acid, NAAG which is one of the best substrates found to date to act as a neurotransmitter in the central nervous system and its abnormal function may be associated with neurotoxic disorder such as epilepsy, ALS, alzheimers etc. PSM carboxypeptidase may serve to process neuropeptide transmitters in the prostate. Neuropeptide transmitters are associated with the

neuroendocrine cells of the prostate and neuroendocrine cells and are thought to play a role in prostatic tumor progression. Interestingly PSM antigen's expression is upregulated in cancer. Peptides known to act as prostatic growth factors such as TGF-a and bFGF, up regulate the expression of the antigen. TNF on the other hand downregulate PSM. TGF and FGF act through the mitogen activated signaling pathway, while TNF acts through the stress activated protein kinase pathway. Thus modulation of PSM expression is useful for enhancing therapy.

#### EXAMPLE 14:

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# 15 IDENTIFICATION OF A MEMBRANE-BOUND PTEROYLPOLYGAMMA-GLUTAMYL CARBOXYPEPTIDASE (FOLATE HYDROLASE) THAT IS EXPRESSED IN HUMAN PROSTATIC CARCINOMA

PSM may have activities both as a folate hydrolase and For the cytotoxic drug a carboxyneuropeptidase. methotrexate to be a tumor toxin it has to get into the cell and be polygammaglutamated which to be active, because polyglutamated forms serve as the enzyme substrates and because polyglutamated forms or toxins are also retained by the cell. Folate hydrolase is a competing reaction and deglutamates methotrexate which then can diffuse back out of the cell. Cells that overexpose folate hydrolase activity are resistant to Prostate cancer has always methotrexate. absolutely refractory to methotrexate therapy and this may explain why, since the prostate and prostate cancer has a lot of folate hydolase activity. However, based prodrugs may be generated which on this activity, would be activate at the site of the tumor such as Nphosphonoacetyl-l-aspartate-glutamate. PALglu is an inhibitor of the enzyme activity with NAAG as a substrate.

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Prostate specific membrane antigen was immuno precipitated from the prostate cancer cell line LNCaP and demonstrated it to be rich in folate hydolase activity, with gammaglutamated folate or polyglutamated methotrexate being much more potent inhibitors of the neuropeptidase activity than was quisqualate, which was the most potent inhibitor reported up to this time and consistent with the notion that polyglutamated folates may be the preferred substrate.

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Penta-gammaglutamyl-folate is a very potent inhibitor of activity (inhibition of the activity of the enzyme is with 0.5um Ki.) As penta-gammaglutamyl-folate may also be a substrate and as folates have to be depolygammaglutamated in order to be transported into the cell, this suggest that this enzyme may also play a role in folate metabolism. Folate is necessary for the support of cell function and growth and thus this enzyme may serve to modulate folate access to the prostate and prostate tumor. The other area where PSM is expressed is in the small intestine. It turns out that a key enzyme of the small intestine that is involved in folate uptake acts as gammacarboxypeptidase in sequentially proteolytically removing the terminal gammaglutaminyl group from In the bone there is a high level of unusual gammaglutamate modified proteins in which the gamma glutamyl group is further carboxylated to produce gammacarboxyglutamate, or GLA. One such protein is osteonectin.

Using capillary electrophoresisis pteroyl poly-gammaglutamate carboxypeptidase (hydrolase) activity was investigated in membrane preparations from androgensensitive human prostatic carcinoma cells (LNCaP). The enzyme immunologically cross-reacts with a derivative of an anti-prostate monoclonal antibody (7E11-C5) that recognizes prostate specific membrane (PSM) antigen. The PSM enzyme hydrolyzes gamma-glutamyl linkages and is an exopeptidase as it liberates progressively glutamates from methotrexate triuglutamate (MTXGlu<sub>3</sub>) and folate pentaglutamate (Pte Glu,) with accumulation of MTX and Pte Glu respectively. The semi-purified membrane-bound enzyme has a broad activity from pH 2 to 10 and is maximally active at pH4.0. Enzymatic activity was weakly inhibited by dithfothreitol ( $\geq 0.2$  mM) but not by reduced glutathione, homocysteine, hydroxymercuribenzoate (0.05-0.5 mM). By contrast to LNCaP cell membranes, membranes isolated from androgeninsensitive human prostate (TSU-Prl, Duke-145, PC-3) and estrogen-sensitive mammary adenocarcinoma (MCF-7) cells do not exhibit comparable hydrolase activity nor a folate hydrolase do they react with 7E11-C5. Thus, LNCap cells that identified in exopeptidase activity and is strongly expressed by these cells.

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PALA-Glutamate 3 was tested for efficacy of the prodrug strategy by preparing N-acetylaspartylglutamate, NAAG 1(Figure 59). NAAG was synthesized from commercially available gamma-benzylaspartate which was acetylated with acetic anhydride in pyridine to afford N-acetylgamma-benzyl aspartate in nearly quantitative yield. The latter was activated as its pentafluorophenyl ester by treatment with pentafluorophenyltrifluoroacetate in pyridine at 0 deg.C for an hour. This activated ester constitutes the central piece in the preparation of compounds 1 and 4 (Figure 60). When 6 is reacted with epsilon-benzyl-L-glutamate in the presence of HOAT(1hydroxy-7-azabenzotriazole) in (tetrahydrofuran, N,N- dimethylformamide) at reflux for an overnight period and after removal of the benzyl protecting groups by hydrogenolysis (H2, 30 psi, 10% Pd/C in ethylacetate) gave a product which was

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identical in all respects to commercially available NAAG (Sigma).

PALA-Glutamate 3 and analog 5, was synthesized in a similar manner with the addition to the introduction of a protected phosphonoacetate moiety instead of a simple acetate. It is compatible with the function of diethylphosphonoacetic acid which allows the removal of the ethyl groups under relatively mild conditions.

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Commercially available diethylphosphonoacetic acid was treated with perfluorophenyl acetate in pyridine at 0 deg.C to room temperature for an hour to afford the corresponding pentafluorophenyl ester in nearly quantitative yield after short path column chromatography. This was then reacted with gammabenzylaspartate and HOAT in tetrahydrofuran for half an hour at reflux temperature to give protected PALA 7 (Nphosphonoacetylaspartate) in 90% yield after flash The free acid was then column chromatography. activated as its pentafluorophenyl ester 8, then it was reacted with delta-benzyl-L-glutamate and HOAT in a mixture of THF-DMF (9:1, v/v) for 12 hours at reflux to give fully protected PALA-Glutamate 9 in 66% yield after column chromatography. Sequential removal of the groups followed by the debenzylation accomplished for a one step deprotection of both the benzyl and ethyl groups. Hence protected PALA-Glutamate was heated up to reflux in neat trimethylsilylchloride for an overnight period. The resulting bistrimethylsilylphosphonate ester 10 was submitted without purification to hydrogenolysis (H, 30 psi, 10% Pd/C, ethylacetate). The desired material 3 was isolated after purification by reverse phase column chromatography and ion exchange resin.

Analogs 4 and 5 were synthesized by preparation of

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phosphonoglutamate 14 from the alpha-carboxyl-protected glutamate.

Commercially available alpha-benzyl-N-Boc-L-glutamate with neat refluxing THF treated at 11 complex to afford the boranedimethylsulfide 90% yield. corresponding alcohol in transformed into bromide 12 by the usual procedure (Pph, CBr<sub>4</sub>).

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The Michaelis-Arbuzov reaction using triethylphosphite to give the corresponding diethylphosphonate 13 which nitrogen with the deprotected at be would trifluoroacetic acid to give free amine 14. The latter with either separately condensed would pentafluorophenylesters 6 or 8 to give 16 and 15 under conditions similar respectively, described for 3. 15 and 16 would be deprotected in the same manner as for 3 to yield desired analogs 4 and 5.

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An inhibitor of the metabolism of purines and pyrimidine like DON (6-diazo-5-oxo-norleucine) or its aspartate-like 17, and glutamate-like 18 analogs would be added to the series of substrates.

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Analog 20 is transformed into compound 17 by treatment with oxalyl chloride followed by diazomethane and deprotection under known conditions to afford the desired analogs. In addition, azotomycin is active only after in vivo conversion to DON which will be released after action of PSM on analogs 17, 18, and 19.

In addition, most if not all chemotherapies rely on one hypothesis; fast growing cells possess a far higher appetite for nutrients than normal cells. Hence, they uptake most of the chemotherapeutic drugs in their proximity. This is why chemotherapy is associated with

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serious secondary effects (weakening of the immune system, loss of hair, ...) that sometimes put the patient's life in danger. A selective and effective drug that cures where it should without damaging what it shouldn't damage is embodied in representative structures 21 and 22.

Representative compounds, 21 and 22, were designed based on some of the specific effects and properties of PSM, and the unique features of some newly discovered 10 cytotoxic molecules with now known mode of action. latter, referred to commonly as enedignes, dynemycin A 23 and or its active analogs. The recent isolation of new natural products like Dynemycin A 23, has generated a tremendous and rapidly growing interest 15 in the medical and chemical sciences. They have displayed cytotoxicities to many cancer cell lines at the sub-nanomolar level. One problem is they are very toxic, unstable, and non-selective. Although they have 20 been demonstrated, in vitro, to exert their activity through DNA damage by a radical mechanism as described below, their high level of toxicity might imply that they should be able to equally damage anything in their path, from proteins to enzymes, ...etc.

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These molecules possess unusual structural features that provide them with exceptional reactivities. Dynemycin A 23 is relatively stable until the anthraquinone moiety is bioreduced hydroanthraquinone 24. This triggers a chain of events by which a diradical species 25 is generated as a result of a Bergman cycloaromatization. species 25 is the ultimate damaging edge of dynemycin It subtracts 2(two) protons from any neighboring molecule or molecules(ie. DNA) producing radicals These radicals in turn combine with molecular therein. oxygen to give hydroperoxide intermediates that, in the 5

case of DNA, lead to single and double strand incision, and consequent cell death. Another interesting feature was provided by the extensive work of many organic chemists who not only achieved the total synthesis of (+)-dynemycin A 23 and other enedignes. but also designed and efficiently prepared simpler yet as active analogs like 26.

Enediyne 26 is also triggerable and acts by virtue of the same mechanism as for 23. This aspect is very relevant to the present proposed study in that 27 (a very close analog of 26) is connected to NAAG such that the NAAG-27 molecule, 21, would be inert anywhere in the body (blood, organs, normal prostate cells, ...etc.) except in the vicinity of prostate cancer, and metastatic cells. In this connection NAAG plays a multiple role:

- Solubilization and transport: analogs of 26type are hydrophobic and insoluble in aqueous media,
  but with a water soluble dipeptide that is indigenous
  to the body, substrate 21 should follow the ways by
  which NAAG is transported and stored in the body.
- 25 Recognition, guidance, and selectivity: Homologs of PSM are located in the small intestines and in the brain.

In the latter, a compound like 27 when attached to a multiply charged dipeptide like NAAG, has no chance of crossing the blood brain barrier. In the former case, PSM homolog concentration in the small intestines is very low compared to that of PSM in prostrate cancer cells. In addition, one could enhance the selectivity of delivery of the prodrug by local injection in the prostate. Another image of this strategy could be formulated as follows. If prostate cancer were a war

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in which one needed a "smart bomb" to minimize the damage within the peaceful surroundings of the war zone, then 21 would be that "smart bomb". NAAG would be its guidance system, PSM would be the trigger, and 27 would be the warhead.

26 and its analogs are established active molecules that portray the activity of dynemycin A. Their syntheses are described in the literature. The total synthesis of optically active 27 has been described. The synthetic scheme that for the preparation of 28 is almost the same as that of 27. However, they differ only at the position of the methoxy group which is meta to the nitrogen in the case of 28. This requires an intermediate of type 29, and this is going to be prepared by modification of the Myers' method. Compound 28 is perhaps the closest optically active analog that resembles very much 26, and since the activity of the latter is known and very high.

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Since NAAG is optically pure, its combination with racemic material sometimes complicates purification of intermediates. In addition, to be able to modify the components of this system one at a time, optically pure intermediates of the type 21 and 22 are prepared. 27 was prepared in 17 steps starting fro commercially available material. Another interesting feature of 27 is as demonstrates in a very close analog 26, it possesses two(2) triggers as shown by the arrows.

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The oxygen and the nitrogen can both engender the Bergman cycloaromatization and hence the desired damage. The simple protection deprotection manipulation of either functionality should permit the selective positioning of NAAG at the nitrogen or at the oxygen centers. PSM should recognize the NAAG portion of 21 or 22, then it would remove the glutamic acid

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moiety. This leaves 27 attached to N-acetylaspartate.

Intramolecular assisted hydrolysis of systems like N-acetylaspartyle is well documented in the literature. The aminoacid portion should facilitate the hydrolysis of such a linkage. In the event this would not work when NAAG is placed on the nitrogen, an alternative would be to attach NAAG to the oxygen giving rise to phenolic ester 22 which is per se labile and removable under milder conditions. PSM specific substrates can be designed that could activate pro-drugs at the site of prostatic tumor cells to kill those cells. PSM specific substrates may be used in treatment of benign prostatic hyperplasia.

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#### EXAMPLE 15:

## GENOMIC ORGANIZATION OF PSM EXON/INTRON JUNCTION SEQUENCES

EXON 1 Intron 1

1F. strand

CGGCTTCCTCTTCGG

10 cggcttcctcttcgg taggggggcgcctcgcggag...tatttttca

1R. strand ...ataaaaagtCCCACCAAA

15 Exon 2 Intron 2

2F. strand

**ACATCAAGAAGTTCT** 

acatcaagaagttct caagtaagtccatactcgaag...

20 2R. strand ...caagtggtcATTAAAATG

Exon 3 Intron 3

3F. strand

25 GAAGATGGAAATGAG

gaagatggaaatgag gtaaaatataaataaataaa...

Exon 4 Intron 4

30 4F. strand

AAGGAATGCCAGAGG

aaggaatgccagagg taaaaacacagtgcaacaaa...

4R. strand ...agagttgTCCCGCTAGAT

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Intron 5 Exon 5 5F. strand CAGAGGAAATAAGGT cagaggaaataaggt aggtaaaaattatctcttttt... ...gtgttttctAGGTTAAAAATG 5 ...cacttttgaTCCAATTT 5R. strand Intron 6 Exon 6 10 6F. strand GTTACCCAGCAAATG gtgaatgatcaatccttgaat... gttacccagcaatg ...aaaaaaagtCTTATACGAATA 6R. strand 15 Intron 7 Exon 7 7F. strand ACAGAAGCTCCTAGA 20 gtaagtttgtaagaaaccargg... acagaagctcctaga ...aaacacaggttatcTTTTACCCA 7R. strand Intron 8 Exon 8 25 8F. strand AAACTTTTCTACACA gttaagagactatataaatttta... aaacttttctacaca ....aaacgtaatcaTTTTCAGTTCTAC 8R. strand 30 Intron 9 Exon 9 9F. strand AGCAGTGGAACCAG agcagtggaaccag gtaaaggaatcgtttgctagca... 35 ...tttctagatAGATATGTCATTC

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9R. strand

...aaagaTCTGTCTATACAGTAA

Exon 10

Intron 10

10F. Strand

CTGAAAAAGGAAGG 5

ctgaaaaaggaagg taatacaaacaaatagcaagaa...

Exon 11 Intron 11

11F. Strand 10

TGAGTGGGCAGAGG

agagg ttagttggtaatttgctataatata...

Exon 13 Intron 12 15

12R. strand

GAGTGTAGTTTCCT

gtagtttcct gaaaaataagaaaagaatagat...

20

Exon 14 Intron 13

13R. strand

AGGGCTTTTCAGCT

agggcttttcagct acacaaattaaaagaaaaaaag...

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Exon 14 Intron 14

14F. strand

GTGGCATGCCCAGG

gtggcatgcccagg taaataaatgaatgaagtttcca... 30

> Exon 16 Intron 15

15R. strand

AATTTGTTTGTTTCC

aatttgtttgtttcc tacagaaaaaaaaaaaaaaa... 35

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Exon 16 Intron 16

16F. strand

CAGTGTATCATTTG

cagtgtatcatttg gtatgttacccttcctttttcaaatt...

5 ...tttcagATTCACTTTTT

16R. strand ...aaagtcTAAGTGAAAA

10 Exon 17 Intron 17

17F. strand

TTTGACAAAAGCAA

tttgacaaaagcaa gtatgttctacatatatgtgcatat...

15 17R. strand ...aaagagtcGGGTTA

Exon 18 Intron 18

18F. strand

20 GGCCTTTTATAGG ggcctttttatagg taaganaagaaaatatgactcct...

18R. strand ...aatagttgTGTAAACCC

Exon 19 Intron 19

19F. strand

GAATATTATATA

gaatattatatata gttatgtgagtgtttatatatgtgtgt...

Notes: F: Forward strand
R: Reverse strand

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#### What is claimed is:

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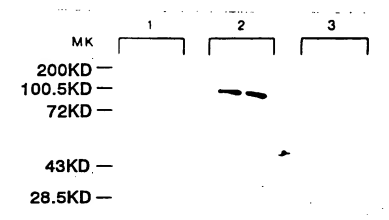
20

- An isolated nucleic acid molecule encoding an alternatively spliced prostate-specific membrane (PSM') antigen.
  - 2. An isolated mammalian DNA molecule of claim 1.
  - 3. An isolated mammalian cDNA molecule of claim 2.
- 4. An isolated mammalian RNA molecule derived from claim 1.
- 5. An isolated nucleic acid molecule of at least 15 nucleotides capable of specifically hybridizing with a sequence of the isolated nucleic acid molecule of claim 1.
  - 6. A DNA molecule of claim 5.
- 7. A RNA molecule of claim 5.
- 8. method of detecting expression of alternatively spliced prostate-specific membrane antigen in a cell which comprises 25 obtaining total mRNA from the cell, contacting the mRNA so obtained with a labelled nucleic acid molecule of claim 5 under hybridizing conditions, determining the presence of mRNA hybridized to thereby detecting molecule, and 30 expression of the alternatively spliced prostatespecific membrane (PSM') antigen in the cell.
- An isolated nucleic acid molecule of claim 2
   operatively linked to a promoter of RNA transcription.

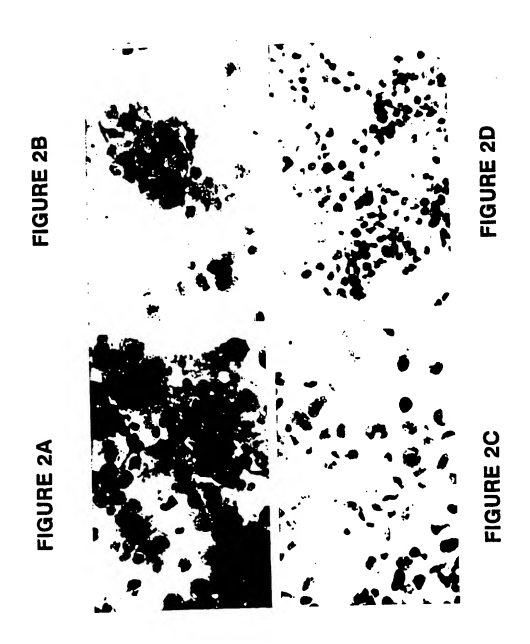
- 10. A vector which comprises the isolated nucleic acid molecule of claim 1.
- 11. A host vector system for the production of a polypeptide having the biological activity of the alternatively spliced prostate-specific membrane (PSM') antigen which comprises the vector of claim 10 and a suitable host.
- 10 12. A host vector system of claim 11, wherein the suitable host is a bacterial cell, insect cell, or mammalian cell.
- 13. A method of producing a polypeptide having the biological activity of the prostate-specific membrane antigen which comprises growing the host cells of the host vector system of claim 12 under suitable conditions permitting production of the polypeptide and recovering the polypeptide so produced.
  - 14. An isolated nucleic acid molecule encoding a prostate-specific membrane antigen promoter.
  - 25 15. A polypeptide encoded by the isolated nucleic acid molecule of claim 1.
  - A method of detecting hematogenous micrometastic 16. tumor cells of a subject, comprising performing nested polymerase chain reaction (PCR) 30 on blood, bone marrow or lymph node samples of the subject using the prostate specific membrane verifying (B) and primers, antigen micrometastases by DNA sequencing and Southern hematogenous detecting thereby analysis, 35 micrometastic tumor cells of the subject.

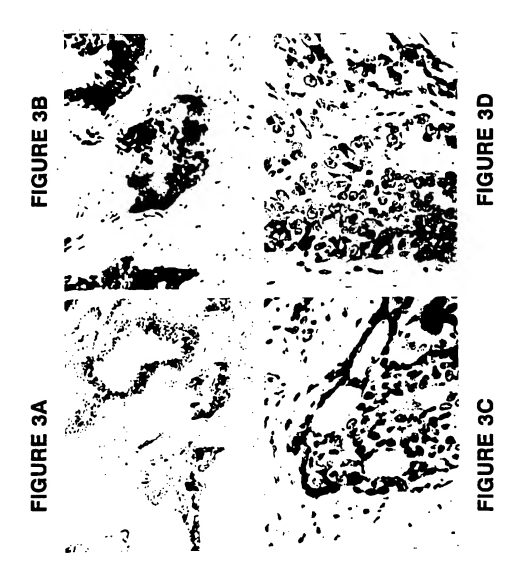
- 17. The method of claim 16, wherein the primers are derived from prostate specific antigen.
- 18. The method of claim 16, wherein the subjects is administered hormones, epidermal growth factor, b-fibroblast growth factors, or tumor necrosis factor.
- prostate cancer of determining method 19. progression in a subject which comprises: a) 10 obtaining a suitable prostate tissue sample; b) extracting RNA from the prostate tissue sample; c) performing a RNAse protection assay on the RNA, thereby forming a duplex RNA-RNA hybrid; d) detecting PSM and PSM' amounts in the tissue 15 sample; e) calculating a PSM/PSM' tumor index, thereby determining prostate cancer progression in the subject.
- 20 20. The method of claim 19, further comprising performing in-situ hyribridization.

#### FIGURE 1



1 - anti- EGFr PoAB RK-2 2 - Cyt-356 MoAB/RAM 3 - RAM





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#### FIGURE 4

100.5

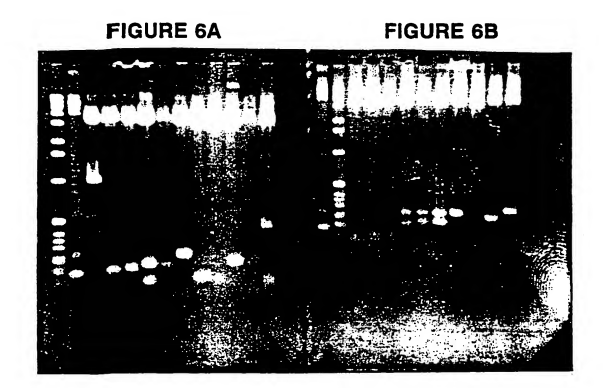
72.0

43.0

28.5

FIGURE 5





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#### FIGURE 7

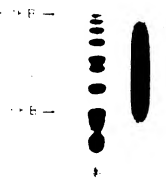
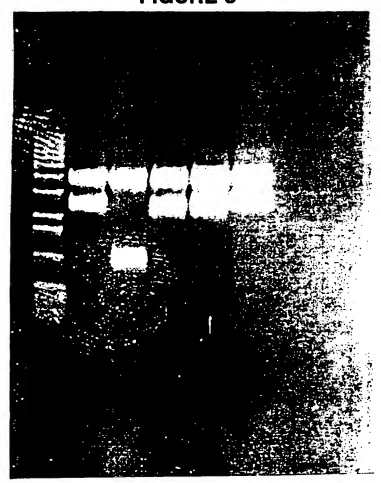


FIGURE 8



#### FIGURE 9

2-1.6-

FIGURE 10

#### FIGURE 11

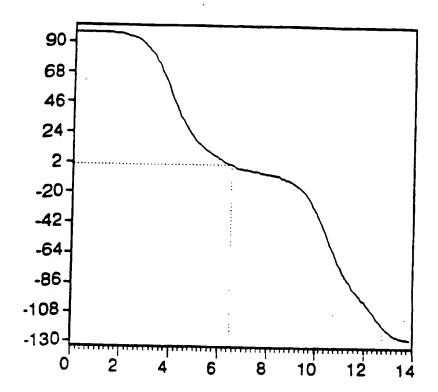
1 2 3

4.4\_\_

FIGURE 12A

FIGURE 12B

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II

II

二

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U

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E

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 $\Xi$ 

田

H

H

国

回

ы

II

II

IX

II

II

IX

II

61

## FIGURE 14-1

on the complete sequence. 750. Total number of residues is: on sequence PMSANTIGEN. Analysis done Done

41.2\$ 10.1% 35.2% **^**|| \ II ۸ II î Z ¥ Z 309 9/ 101 CNAT CNAT CNAT CNAT -75 -88 0 11 II Π П 20] 20 conformation conformation conformation conformation E E E C Extended Helical Turn Coil In In In

14/130

conformation codes. Sequence shown with

are given conformation Ø in more residues or S of stretch Consecutive overlined.

161 II E II 15 II 回 IX 161 II IM II IE II H 1= H II 回 II 163 IX H H 10 臼 10 10 ы 回 10 II IU II 10 II II II II II IH IX II II II II IX II II 161 IX IX E 二 IM IX 田 II II 回 IX 工 回 II 31

# FIGURE 14-1

sednence. 750. Analysis done on the complete residues is: on sequence PMSANTIGEN. Total number of Done

35.2\$ 10.1% î **^**|| **^**|| î ¥ Z 309 92 264 101 CNAT CNAT CNAT CNAT -75 -88 0 n n n 20 20] conformation conformation conformation conformation E) (E) Extended Helical Turn Co11

14/130

conformation codes. Sequence shown with

given conformation Ø in more residues or വ of stretch Consecutive overlined

are

16 II II 回 IX 二 H II I II I 田 IX 10 回 IEI II 10 E II 10 E 1= 10 H IX 10 12 II II 回 H 10 F H 10 国 H M 10 H 10 E 工 10 10 II IX 王 二 II II 工 IX II II IX II I II IX II IX IX II II H II IX E II II 回 II II II 田 IX IE IX IX H II 31 61

[L]

E

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E

E

回

FIGURE 14-3

H	E	H	II	II	IEI	回	lEI	Ö	IX
H	मि	Ö	II	II	E	IE	i E	U	IX
I		M	IX	IX	IEI	IEI	लि	II	II
X	ास	H	IX	H	E	ပ	IEI	IX	II
X	IEI	H	IX	IEI	E	ပ	E	II	II
X	H	E	II	लि	E	H	164	II	IX
H	10	দ্র	ΙΞ	IEI	ाध	H	回	IΞ	IX
I	10	(L)	IX	F	II	E	H	IX	Œ
Œ	10	O,	IX	लि	II	II	ပ	II	IX
E)	IO	Ö	II	नि	II	ΙΞ	ပ	II	I
[E]	IO	ပ	IX	II	IX	I	ပ	I	II
E	H	E	I	II	IX	ΙΞ	H	E	ΙΞ
(L)	H	I	IX	II	U	IX	E	E	IX
(±)	F	I	II	II	ບ	IX	E	H	E
Œ	IΞ	II	धि	IX	X	IX	E	回	
F	IX	II	लि	ाध	नि	IX	王	लि	E
Œ	II	ΙΞ	IEI	ļШ	E	IX	I	<b>E</b>	नि
Œ	II	E	ाध	नि	E	II	IH	E	<b>L</b>
Œ	II	E	नि	।ध्य	E	ΙX	I	िल	E
F	IX	E	नि	E	E	II	Œ	ाध	IЫ
E	IX	U	नि	IEI	II	IX	I	H	IEI
Ħ	IX	U	lm	lΠ	IΞ	IX	II	ပ	I
H	II	U	U	EI	ΙΞ	II	I	ပ	IX
E)	I	D	IH	lei I	II	H	IX	ပ	I
F	H	10	Œ	Ξ	IX	IX	II	U	II
U	II	lei	IE	ल	IX	IX	II	H	II
ပ	I	E	IH	II	IX	IX	IX	<b> </b>	IX
团	U	ि	IE	IΞ	IX	IX	E	IM	IX
E)	ບ	IEI	E	II	II	IX	E	IM	IX
田	I	E	H	I	IX	H	IП	iei	ບ
451	481	511	541	571	601	631	661	691	721

FIGURE 14-4

Semi-graphical output.

Symbols used in the semi-graphical representation:

conformation: conformation: Extended **Co11** conformation: conformation: Helical Turn

50 **MWNLLHETDSAVATARRPRWLCAGALVLAGGFFLLGFLFGWFIKSSNEAT** 30 XXXXXXXXXXXXXX 20

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100 X<\*\*\*\*\*\*\*\*\*\* 80 XXXXXXXXXXXXX 9

X<\*\*\*\*\*\*XXXXX----

nitprhnmkafldelkaenikkflynftqiphlagteqnfqlakqiqsqw

FIGURE 14-5

XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
XX - X - X - X - X - X - X - X - X - X

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350  ****  ****  ****
GNFSTQKVKM *-*XXXXXX- *-*XXXXXX- 390 GIDPQSGAAV
VPYNVGPGFT
PPDSSWRGSLK **>->>-> **>->>->
GSA:

FIGURE 14-7

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XXX	450 	X	500      ESWTKK	(XX>>>*	550    KFSGYP	1-4<<	009
XXX##<###<<	440   Waeensrllq	-*XXXXXXXXX*****XXXXXXXXXX-	490    PDEGFEGKSL)	-XXXXXXXXX**XXXXXXXXXXXXXXXX-	540   Varytknweta	-+<<<<<-<	965 
	430   EEFGLLGSTE	XXXXX	480   VHNLTKELKS]	XXXXXX**) XXXXXXX**)	530     FFQRLGIASGE	CXXXX>+++	280
	420   RTILFASWDA		470   VDCTPLMYSL	(XX	520     Klgsgndfevi	(	570
	410 420 430 440 450           SFGTLKKEGWRPRRTILFASWDAEEFGLLGSTEWAEENSRLLQERGVAYI	XXX***>>>****>- XXX***>>>****>-	460 470 480 490 500           NADSSIEGNYTLRVDCTPLMYSLVHNLTKELKSPDEGFEGKSLYESWTKK		510 520 530 540 550           SPSPEFSGMPRISKLGSGNDFEVFFQRLGIASGRARYTKNWETNKFSGYP		560

FIGURE 14-8

LYHSVYETYELVEKFYDPMFKYHLTVAQVRGGMVFELANSIVLPFDCRDY	XXX <xxxxxx< th=""><th>XXXXXXXXX-XXXXXXXXXXXXXXXX</th></xxxxxx<>	XXXXXXXXX-XXXXXXXXXXXXXXXX
--	--	----------------------------

XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	620 630 640 650	AVVLRKYADKIYSISMKHPQEMKTYSVSFDSLFSAVKNFTEIASKFSERL	XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX
		I KIYSISMKHPQEMKTY	XXXXX**XX
	610	AVVLRKYADI	XXXXXXXX

XXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXXX	700	(APSSHNKY
	069	, PDRPFYRHVIY
(XX)	089	RAFIDPLGLE
XXX**X	670	WMNDQLMCLE
XXXXXXXXXXXXXXXXXXXXXXXXXXXX	099	QDFDKSNPIVLRMMNDQLMCLERAFIDPLGLPDRPFYRHVIYAPSSHNKY

<b>*</b> *	<b>* 4 4</b>
****	44 4
1	i   
<u>.</u>	1
<	^
XXXX>>	KXXX>>
XXX	XXX
XXX	XXX
XX	XXX
×-	-XX
į	į
i	i .
>>*********	(X##<<<
×××	XX

750	_	<b>AETLSEVA</b>
740	_	FTVQAA
730		<b>AWGEVKRQIYVAA</b>
720		IESKVDPSKA
710		<b>AGESFPGIYDALFDIESKVDPSKA</b>

----XXXXXXX\*\*\*\*XXXXXXXX ---XXXXXXX\*\*\*\*XXXXXXX--<--

22/130 FIGURE 15A

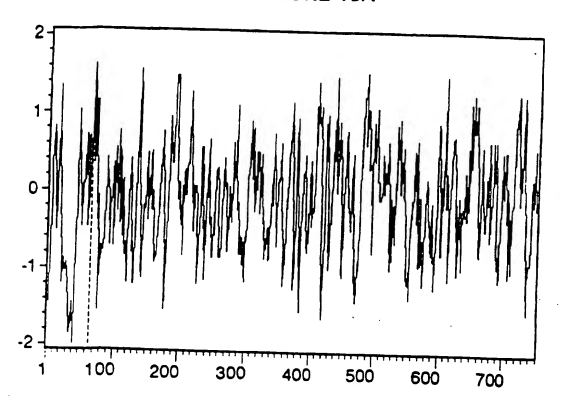


FIGURE 15B

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The three highest points of hydrophilicity are:

-> This is the value recommended by the authors

The averaging group length is: 6 amino acids.

The method used is that of Hopp and Woods.

the complete sequence.

Total number of residues is: Done on sequence PMSANTIGEN.

Analysis done on

ANTIGENIC DETERMINANTS

· PREDICTION OF

\*

Asp-Glu-Leu-Lys-Ala-Glu **89** 63 From

Asn-Glu-Asp-Gly-Asn-Glu Lys-Ser-Pro-Asp-Glu-Gly 137 487 to to 32 482 From From 1.57 1.55 7

hydrophilicity. Average for: Ah stands

only the highest point was in 100% The second and third points group. of incorrect predictions control proteins, a known antigenic to Note that, on a group of proportion of 33% of the cases assigned gave a

24/13	30
-------	----

RATTRFR Rat transferrin receptor mRNA, 3' end. 164  HUMTFRR Human transferrin receptor mRNA, complete cd 145  CHKTFER G.gallus mRNA for transferrin receptor 203  51.9% identity in 717 nt overlap  1020 1030 1040 1050 1060 107  pmsgen TGTCCAGCGTGGAATATCCTAAATCTGAATGGTGAGACCCTCTCACAC  CHKTFE TACACTTATCCCATTCGACATGCCCACTTGGAACTGGAGACCCTTACACC  990 1000 1100 1110 1120 1130  pmsgen CCCAGCAATAGCTTATAGGCGTGGAATTGCAGAGGCTGTTGGTCTTCC  1080 1090 1130  CHKTFE CCCTTCGTTCAACCACACCCAGTTTCCACACTTCAGGACTACC  1050 1050 1060 1070 1080	RATTRER Rat transferrin receptor mRNA, 3' end. 164 HUMTFRR Human transferrin receptor mRNA, complete cd 145 HUMTFRR Human transferrin receptor 203 51.9% identity in 717 nt overlap 1020 1030 1040 1050 1060 107 pmsgen TGTCCAGCGTGGAATATCCTAAATCTGAATGGTGCAGAGCCCTTACACC 990 1000 1100 1110 1120 1130 pmsgen CCCAGCAATGATGCTTATAGGCGTGGAATTGCAGGGCTGTTGGTTTC :::::::::::::::::::::::::	The best CHKTFER	FIGURE 16-1 Scores are: G.gallus mRNA for transferrin receptor	initn 201		initi
CHKTFER G.gallus mRNA for transferrin receptor 203  51.9% identity in 717 nt overlap  1020 1030 1040 1050 1060 107  pmsgen TGTCCAGCGTGAAATATCTGAATGGTGCAGAGACCCTTCACACCCCG990 1090 1100 11020 1030  1080 1090 1100 1110 1120 1130  pmsgen CCCAGCAATGATTATAGGCGTGGAATTGCAGAGGCTGTTGGTTTGGTTTTCCCCCG990 1130  CHKTFE CCCTTCGTTCAACACCACACCCACCTTTCGTTTTTCCTTTCTTT	CHKTFER G.gallus mRNA for transferrin receptor 203 51.9% identity in 717 nt overlap 1020 1030 1040 1050 1060 1070 pmsgen TGTCCAGCGTGGAATATCCTAAATCTGAATGGTGCAGGAGCCTTCACACCGGAGACTTACCCATTCGACTGCACTGGAACTGGAACTGCAGACCCTTACACCCCGGAATTCCAATTCCCATTCGAATTGCTAATTGAATTGCAAATTGAAATTGAATTGAAAAAAAA	RATTRE		164		164
CHKTFER G.gallus mRNA for transferrin receptor 203 51.9% identity in 717 nt overlap 1020 1030 1040 1050 1060 1070 pmsgen TGTCCAGCGTGGAAATATCCTAAATCTGAATGGTGCAGGAGACCCTCTCACACC 990 1000 1000 1110 1120 1030 1130 pmsgen CCCAGCAAATGATATGCTTATAGGCGTGGAATTGCAGAGGCTGTTGGTTTCC 990 1090 1100 1110 1120 1130 cCCAGCAAATGAATATGCTTATAGGCGTGGAATTGCAGAGGCTGTTGGTTTCC ii: i:	CHKTFER G.gallus mRNA for transferrin receptor 203 51.9% identity in 717 nt overlap 1020 1030 1040 1050 1060 1070 pmsgen TGTCCAGCGTGGAATTATCCTAAATCTGAATGGTGCAGGAGCCTTCACACC 990 1000 1010 1120 1130 pmsgen CCCAGCAATGCTTATAGGCGTGGAATTGCAGAGCTGTTCCC  CHKTFE CCCTTCGTTCAACCACCCCACTTCAGAATTGCTTTCCCCCCAGTTTCCCCCCAGTTTCCTTCC	HUMTFR	Human transferrin receptor mRNA, complete	145		145
	1020	CHKTFE 51.9	Įd	203	• •	120 321
		pmsgen	1020 1030 1040 1050 1060 FGTCCAGCGTGGAATATCCTAAATCTGAATGGTGCAGGAGACCCTCT	107 CACAC	ం చ	GGTT
1080 1090 1100 111 CCCAGCAAATGAATATGCTTATAGGCGTGGAA1 ::: : : : : : : : : : : : : : : : : :		CHKTFE		CACCC	CA 10	::::: CAGGCT' 1040
		pmsgen CHKTFE		113 TCTTC ::: ACTAC	O C C C C	AGTAT :: CACAT
	TGCTGTTCAGACCATCTCTAGCAGTGCAGCAGCCAGGCTGTTCAGC	bmsgen		AATGG	3TG	GCT(
				•••	••	••
				AATGG	√T'G(	GAG

pmsgen CHKTFE	AGCACCACCAGA :: : : : : : : : : : : : : : : : : : :	1210 CAGATAGCAG : :: :: ; CTGA-AGC	10 1220 TAGCAGCTGGAGGAA :: :::::::: -AGGTTGGAAAGGTG	1230 AAGTCTCAAAGT ::: TGCGATCCA	1240 12 GCCCTACAATGTT ::: :: :: TTCCTGTAAGGT-	pmsgen AGCACCACCAGATAGCAGCTGGAGGAAGTCTCAAAGTGCCCTACAATGTTGGACCTGG :: :: :: :: :: :: :: :: :: :: :: :: ::
pmsgen CHKTFE	1260 12 pmsgen ctttactggaaa : :::: chktfe caaagcaggaga	1270 SAAACTTTTC ::::::::::::::::::::::::::::::	1280 :TACACAAAAAG : : : : : :CAGA-TAATGG 1230	1280 1290  FACACAAAAGTCAAGATGCAC : :: :: :: :: :: :: :: :: :: :: :: :: :	1300 131 CATCCACTCTACCA : :: ::: TGTGAACAATTCCA 1250 12	1260 1270 1280 1290 1300 1310  CTTTACTGGAAACTTTTCTACACAAAAAGTCAAGATGCACATCCACTCTACCAATGAAGT : :::::::::::::::::::::::::::::::::::
pmsgen	1320 GACAAGAATT : ::: CAGGAAGATT 1270	1330 TTACAATGT : :: : TCTGAACAT	0 1340 1350 13 AATGTGATAGGTACTCTCAGAGGAGCAGT :: ::::::::::::::::::::::::::::::::::	1350 TCAGAGGAGC:::::TCCAGGGATT	1360 AGTGGAACCA : :::: TGAAGAACCT 1310	1320 1330 1340 1350 1360 1370 pmsgen GACAAGAATTTACAATGTGATAGGTACTCTCAGAGGAGCAGTGGAACCAGACAGA
pmsgen (CHKTFE 1	1380 13 CATTCTGGGAGG : :::: TGTGATTGGAGC	1390 AGGTCACCG(:: :: :: :: AGCCCAGAGA	90 1400 1410  TCACCGGGACTCATGGGTGTTTGG :: :::: ::: :::  CCAGAGACTCCTGGGGCCCAGG 340 1350 1360	1410 TGTTTGGTGG' GCCCAGGAGTG	1420 TGGTATTGACCCT : : : :: AGTGGCTAAAGCT 1370	pmsgen CATTCTGGGAGTCACCGGGACTCATGGGTGTTTGGTGGTATTGACCCTCAGAGTGGAGC : :::::::::::::::::::::::::::::::::::

		/130	
1440 1450 1460 1470 1480 1490 pmsgen AGCTGTTGTTCATGAAATTGTGAGG——GAGCTTTGGAACACTGAAAAAGGAAGGTGGAG :::::::::::::::::::::	1500 1510 1520 1530 1540 1550  pmsgen ACCTAGAAGAATTTTGTTTGCAAGCTGGGATGCAGAAGAATTTTGGTCTTCTTGGTTTC  :::::::::::::::::::	TGGCTTATATTAA : ::::::::::::::::::::::::::::	FACACCGCTGATG  SECOCCTTGCTG  CAGCCCCTTGCTG
1480 CACTGAAAAAG : :::::: FAGTGAAAAAC 1430	1540 \AGAATTTGGT( ::::::::::::::::::::::::::::::::	1600 GCGTGGCGTGG : : CAAAGCTTTC/ 1550	GTTGATTGTACACCGCTGA
1470 1480 1490 GAGCTTTGGAACACTGAAAAAGGAAGGGTG :::::::::::::::::::::::	1500 1510 1520 1530 1540 1550  pmsgen ACCTAGAAGAATTTTGTTTGCAAGCTGGGATGCAGAAGAATTTGGTCTTCTTGGTTC  :::::::::::::::::::::	1560 1570 1580 1590 1600 1610  pmsgen TACTGAGTGGGCAGAGTTCAAGACTCCTTCAAGAGCGTGGCGTGGCTTATTTAA  :::::::::::::::::::::::::::::	1620 1630 1640 1650 1660 1670  pmsgen TGC-TGACTCATCTATAGAAGGAAACTA-CACTCTGAGAGTTGATTGTACACCGCTGATG  :::::::::::::::::::::::::::::::::
0 1460 GAAATTGTGAGC :::::::: GAACTTGCCCGTGTG 00 1410	1520 TGTTTGCAAGC: X:::::::TTTGCTAGC	570 1580 AGGAGAATTCAAGA :::::X AGGGGTACTCTGCC 0 1530	0 1640 GAAGGAAACTA :::::: CCTGGGAGCAA
1450 GTTCATGAAA : :: NTGTTGGAAC	1510 AGAACAATTT ::::::	1570 TGGGCAGAGG, ::: ::: TGGCTGGAGG	0 1630 CTCATCTATAG : ::: ATGCTCCAGTC
1440 n AGCTGTT :::: E TGCTATA 1390	1500 :: :: ::: E ACCGAGGCGAAG	1560 n TACTGAGTGGG ::::::: E TACTGAATGGC	1620 n TGC-TGACTC/ :::::: E -GCTTGGATG(
pmsge	pmsge	pmsgen CHKTFE	pmsgen CHKTFE

1680

FIGURE 16-4

pmagen tacagettggtacacaacetaacaaagagetgaaaagecetgatgaaggetttgaagge

TATATGCTGCGGGGGTATTATGAAGGGGGTGAAGAATCCAGCAGTCTCAGAGAGG CHKTFE

1660

1650 1630

--CTCTATAACAGACTTGGCCCAGACTGGGTAAAAGCAGTTGTTCCTCTTGGCCTGGA

CHKTFE

116 104 107	1240 1250 AAAGTGCCCTACAATGTTGGACCTGGCTT- :::::::::::::::::::::::::::::::::::	1310 ACTCT-ACCAATG ::::::: ACTGTGAACAATGTACT 720
RATTRFR Rat transferrin receptor mRNA, 3' end. 55.5% identity in 560 nt overlap	pmsgen ccaccagaragcrggagaggaagrcrcaaagrgcccracartgrrggaccrggcrr- :::::::::::::::::::::::::::::::::::	1260 1270 1280 1290 1300 1310 pmsgen -TACTGGAAACTTTTCTACACAAAAAGTCAAGATGCACATC-CACTCT-ACCAATG RATTRF CTCATGTAAGCTGGAACTTTCACAGAATCAAAATGTGAAGCTCACTGTGAACAATGTACT 670 680 690 700 710

	13	1320	1330	1340		1350	1360	1370
pmagen	AAGTGA	ACAAGAATT	FTACAATC	TGATAGG	TACTOTO	AGAGGAG	pmsgenAAGTGACAAGAATTTACAATGTGATAGGTACTCTCAGAGGAGCAGTGGAACCAGACAG	GACAG
RATTRF	GAAAGAAA	CAAGAATA	CTTAACA	TCTTTGG	CGTTATI	AAAGGCT	RATTRF GAAAGAAACAAGAATACTTAACATCTTTGGCGTTATTAAAGGCTATGAGGAACCAGACCG	GACCG
7.	730	740	750	760	0	770	780	
	13	1380	1390	1400		1410	1420	1430
pmsgen	pmsgen ATATGTCATTCT	TTCTGGGA	GGTCACC	GGGACTC	ATGGGTG	TTTGGTG	GGGAGGTCACCGGGACTCATGGGTGTTTTGGTGTATTGACCCTCAGAG	CAGAG
RATTRF	RATTRF CTACATTGTAGT	TAGTAGGA	GCCCAGA	GAGACGC	TTGGGGC	GGAGCCCAGAGACGCTTGGGGCCCTGGT-GTTGCGA	AGGAGCCCAGAGACGCTTGGGGCCCTGGT-GTTGCGAAGTCCAGTG	CAGTG
75	790	800	810	820	0	830	840	
	7	1440	1450	1460	0	1470	1480	
pmsgen	T-GGAGCA	GCTGTTGT	TCATGA	ATTGTGA	GGAGCTT	TGGAACA-	pmsgen T-GGAGCAGCTGTTGTTCATGAAATTGTGAGGAGCTTTGGAACA-CTGAAAAAGGAA	AGGAA
	••	•••	••	••	••		••	••
RATTRF	TGGGAACA	GGTCTT-C	TGTTGA	ACTTGCC	CAAGTAT	TCTCAGA	RATTRF TGGGAACAGGTCTT-CTGTTGAAACTTGCCCAAGTATTCTCAGATATGATTTCAAAAGAT	AAGAT
~	850	860	870		880	890	006	
	1490	1500	15	1510	1520	1530	) 1540	
pmsgen	GGGTGGAG	ACCTAGAA	GAACAAT	TTTGTTT	GCAAGCT	GGGATGCA	pmsgen GGGTGGAGACCTAGAAGAACAATTTTGTTTGCAAGCTGGGATGCAGAAGAATTTGGTCTT	GTCTT
	:::: X	••	••••••		•••	••	•••••••••••••••••••••••••••••••••••••••	••
RATTRF	GGATTTAG	ACCCAGCA	GGAGTAT	TATCTTT	GCCAGCT	GGACTGCI	RATTRF GGATTTAGACCCAGCAGGAGTATTATCTTTGCCAGCTGGACTGCAGGAGACTATGGAGCT	GAGCT
	910	920	930		940	950	096	

3	30/130	
1600 CGTGGCGTG : : 1020	1660 TGATTGTAC : : : TTCTGCCAG 1080	1720 CTGATGAAG : :: CCGA
1590 CTCCTTCAAGAG : :: :: TTGCATCTAAAG	1650 CTCTGAGAGT : : :: ACTTCAAGGT 1070	1710 TGAAAAGC-C : ::::
1580 1590 TTCAAGACTCCTTCAAGAGC :::: ::: ::: CCTTTCATCTTTGCATCTAAAG- 1000 1010	1640 1650 1 AGGAAACTA-CACTCTGAGAGTTGATT :::::::::::::::::::::::::::	ACAACCTAACAAAGAGCTGAAAAGC-CCTG
0 1570 GTGGCAGAGGAGAA:::: X GTGGCTGGAGGGGTACG	1630 ATCTATAGAAG : : : AAGTCGTCCTG	1690 FACACAACCTA :: ::
1560 CTGAGTGGGC ::::::::	1620 ATGCTGACTC/ :: :: : : AT-CTGGATA/ 1040	1680 [ACAGCTTGG] ::::::::::::::::::::::::::::::::::
pmsgen CTTGGTTCTACTGAGTGGCAGAGAATTCAAGACTCCTTCAAGAGCGTGGCGTG	pmsgen GCTTATATTAATGCTGACTCATCTATAGAAGGAAACTA-CACTCTGAGACTTGATTGTAC ::::::::::::::::::::::::::::::::::::	pmsgen ACCGCTGATGTACAGCTTGGTACAACCTAACAAAAGAGCTGAAAAGC-CCTGATGAAG :::::::::::::::::::::::::::::::
pmsgen RATTRF	pmøgen Rattrf	pmsgen

	1730	1740		1750	1760	1770
pmsgen	pmsgen GCTTTGAAGGCJ	AAATCTCTTT	AT-GAA	AAATCTCTTTAT-GAAAGTTGGACTAAAAAAAAGTCCTTCCCCAG	NAAAAAAGTC	CTTCCCCAG
)	••	•••	•••	••	••	
RATTRF	TTGATGGAAAATATCTATATCGAAACAGTAATTGGATTAGCAAAATTGAGGAACTTT	AAATATCTAT	ATCGAAACAG	TAATTGGATTA	GCAAAATTG	AGGAACTTT
	1140	1150	1160	1170	1180	1190
	1780	1790	1800	1810	1820	1830
pmsgen	pmsgen AGTTCAGTGGCA	ATGCCCAGGAT	LAAGCAAATT	<b>ATGCCCAGGATAAGCAAATTGGGATCTGGAAATGATTTTGAGGTGTTTT</b>	ATGATTTTG	AGGTGTTCT
RATTRE	RATTRF CCTTGGACAATGCTGCATTCCTTTTTTTTTTCAGGAATCCCAGCAGTTTCTTTC	GCTGCATTCCC	TTTTCTTGC	ATATTCAGGAA	TCCCAGCAG	FTTCTTTCT
	1200	1210	1220 1230	1230	1240	1250

266 145 Human transferrin receptor mRNA, complete cd overlap identity in 464 nt HUMTFRR

	32/130	
1230 1240 1250 1260 1270  pmsgen AGGAAGTCTCAAAGTGCCTACAATGTTGGACCTGGCTTTTAC-TGGAAACTTTTTTACAC :::::::::::::::::::::::::	1280 1290 1300 1310 1330 pmsgen AAAAAGTCAAGATGCACTCT-ACCAATGAAGTGACAAGAATTTACAA ::::::::::::::::::::::::::::::::::	pmsgen TGTGATAGGTACTCTCAGAGGAGCAGTGGAACCAGACAGA
TTTC:	AATT	3GGA(
1270 AACTT : GGATG 1190	1320 AAGTGACAAGAATT :::::::: AAGAGATAAAAATT 0	GATATGTCATTCTGGGAGG
GGAA : : GTAG	1320 TGACA :::	1380 TCATT
AC-T :::: ACAT 80	AAG ::: TGAAAG 1240	OGATATG
1260 CTTTAC- :: ::: CTCTACA	 3CTG	1370 ACAGA : ATCAC
1260   12   ACCTGGCTTTAC-TGGAAA   : : : : : : : : : : : : : : : : : : :	\TG \TGT(	1360 CAGTGGAACCAGACA :::::::::
50 GGACC : : AAAAC 1170	1310 ACCAA: ::: AGCAA:	GAACC
1250 GTTGG TGGAA	1300 1310 ATC-CACTCT-ACCAATG : : : : : : : : : : : : : : : : : : :	1360 IGTGG :: :
CAAT TGAC	0 CACI CACI	AGCA
1240 CCCTAC CCCTCT	1300 ATC-C, : : : AGCTC, 1220	1350 CAGAGGAG : ::: TAAAGGCT
GTGC GTGC TGTC	GCAC	13 CTCA : :
30 TCAAA GAGAC	290 AAGATG : ::: AGAATG	O GTACT : : GAGTT
1230 STCTC	1290 STCAAG : : SCAAGA 121	1340 GATAGGT
GAAG	AAAAGTO	TGAT
n AG R TA	1280 en AA : : FR AG	n TG
nsgel JMTF1	1; nsgel jwtf	nsge! JMTF
<u>a</u> <del>x</del>	ā	

pmsgen ccgcgacrcarcgcrcrrcgrcrrcacrccrcacagr-ccaccrcrra HUMTFR GAGAGATGCATGGGGCCCTGGAGCTGCAAAATC-CGGTGTAGGCACAGCTCTCTCTTTGA 1440 1360 1430 1420 1410 1400

pmsgen HUMTFR	- AAATTG :: :: AACTTGCC	1460 -TGAGGAGCTT :::::: CAGATGTTCTC 1390	1470 FTGGAACACTG : : : PAGATATGGTC	1480 1490 GAAAAAGGAAGGGTG( ::: X:: CTTAAAAGATGGGTT 1410 1420	3AG FCA	1500 ACCTAGAAGAACAA :: :: :: :: :: :: :: :: :: :: :: :: ::
pmsgen HUMTFR	pmsgen TTTTGTTTGCAAAUMTFR TTATCTTTGCAAAUMTFR TTATCTTTGCCAAAAAAAAAAAAAAAAAAAAAAAAAAAA	1520 GCAAGCTGGGA :::::: GCCAGTTGGAG 1450	20 1530 1540 AGCTGGGATGCAGAAGAATTTGG :::::::::::::::::::::::::::	1540 TTTGGTCTTC ::::: TTTGGATCGG	pmsgen TTTTGTTTGCAAGCTGGGATGCAGAAGTTTTGGTCTTTTTTTT	1560 TGAGTGGGCAG ::::::: TGAATGGCTAG
pmsgen HUMTFR	1570 A-GGAGAA7 : ::: : AGGGATACC 1500	1580 PTCAAGACTCC : : : :: CTTTCGTC-CC 1510	1590 TTCAAGAGCG7: :: TGCATTTAAAG	1600 FGGCGTGGCT : :: SGCTTTCACT 1530	1570 1580 1590 1600 1610 1620  pmsgen A-GGAGAATTCAAGACTCCTTCAAGAGCGTGGCGTGGCTTATATTAATGCTGACTCATCT :::::::::::::::::::::::::::::::	1620 TGACTCATCT :: :: GGATAAAGCG 1550
pmsgen	1630 ATAGAAGGA : ::	1640 AAACTACACTC	1650 TGAGAGTTGA1 : : :: :	1660 FTGTACACCG	1630       1640       1650       1660       1670       1680         ATAGAAGGAAACTACACTCTGAGGTTGATTGTACACCGCTGATGTACA-GCTTGGT-AC	1680 GCTTGGT-AC::::::::::
HUMTFR	GTTCTTGG1 1560	FACCAGCAACT 1570	TCAAGGTTTC1 1580	TCTGCCAGCCCA	HUMTFR GTTCTTGGTACCAGCAACTTCAAGGTTTCTGCCAGCCCACTGTTGTATACGCTTATTGAG	TACGCTTATTGAG 1610

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1740	Paragan nemectanenandaelenandeelelenandeellisaabseenmerriellinair	HUMTFR AAAACAATGCAAAATGTGAAGCATCCGGTTACTGGGCAATTTCTATATCAGGACAGCAAC	1670
1730	TITCAAGGC	ATTTCTATA	1660
1720	TOWI CAMES	TTACTGGGCA	1650
1710	ופאאאפרכי	AAGCATCCGG	1640
1700	.::::	SCAAAATGTG	1630
1690	***	AAAACAAT	1620
		HUMTFR	

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## 35/130 FIGURE 17A



FIGURE 17B



FIGURE 17C



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FIGURE 18

1 2

100 –

68 –

43 -

FIGURE 19

1 2 3 4

200 kDa — PSI
69 kDa —

#### FIGURE 20

1 2 3 4 5 6 7 8 9 10 11 12 13 14 15

400

350

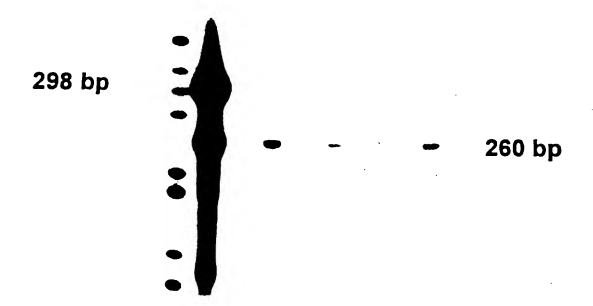
FIGURE 21

1 2 3 4 5 6 7 8 9 10

298 bp 260 bp

FIGURE 22

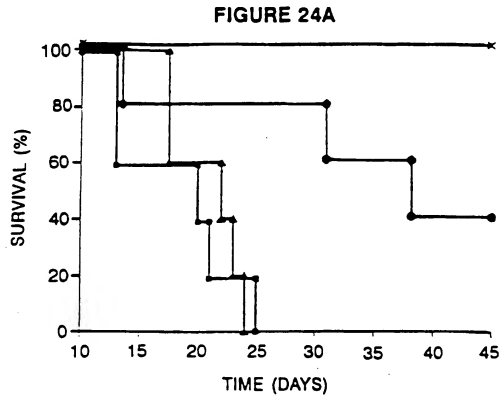
1 2 3 4 5 6 7 8 9



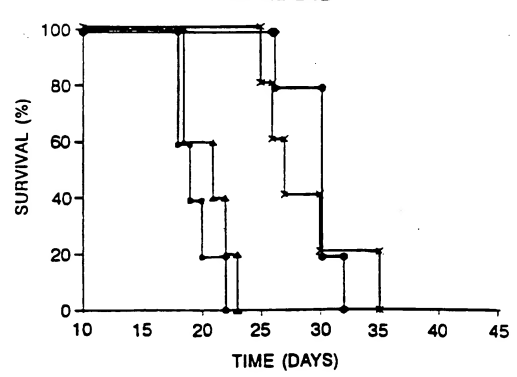
#### 41/130 FIGURE 23

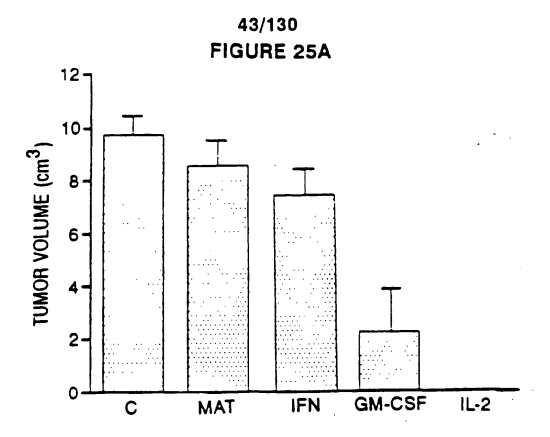
CELL LINE/TYPE	11p11.2-13 REGION	METASTATIC	PSM RNA DETECTED	PSM DNA DETECTED
LNCap			++	ND
HUMAN PROSTATE		·	++	ND
A9 (FIBROSARCOMA)	NO	NO	-	-
A9(11) (A9+HUM. 11)	YES	NO	_	REPEAT
AT6.1 (RAT PROSTATE)	NO	YES	_	-
AT6.1-11-c11	YES	NO	+	++
AT6.1-11-c12	NO	YES	-	_
R1564 (RAT MAMMARY)	NO	YES	-	-
R1564-11-c14	YES	YES	-	+
R1564-11-c15	YES	YES	_	REPEAT
R1564-11-c16	YES	YES	-	ND
R1564-11-c12	YES	YES	ND	+

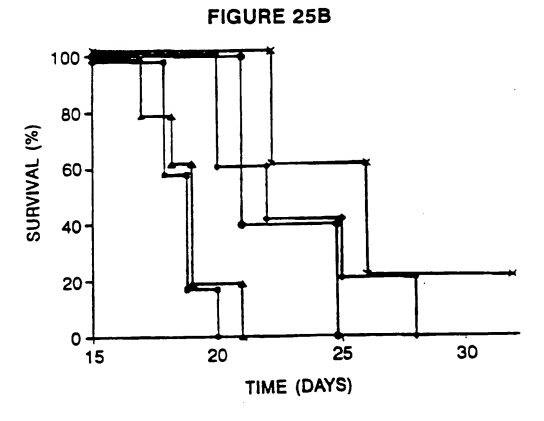
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#### FIGURE 26

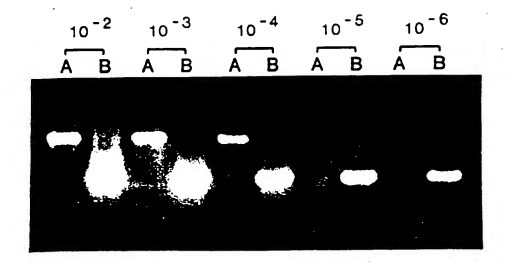
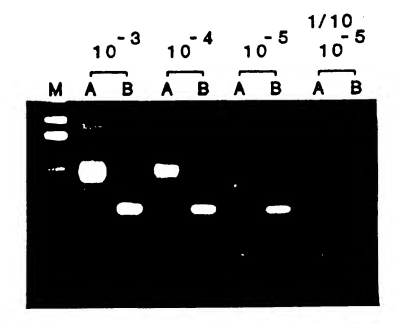
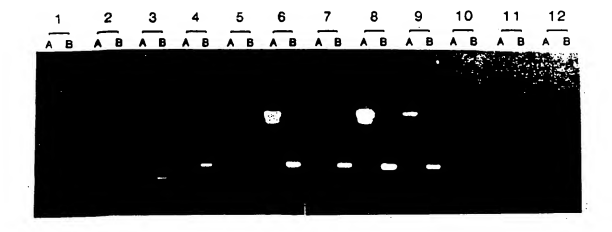


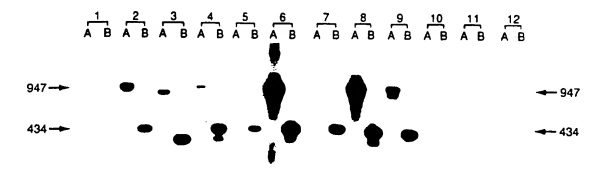
FIGURE 27



## FIGURE 28



#### FIGURE 29



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			_	
		UR		30
Г	u	JN	_	JU

Patient	Stage	Treatment	PSA	PAP	PSA-PCR	PSM-PCR
1	T2NxMo	None	8.9	0.7	_	+
2	T2NoMo	RRP 7/93	6.1	_	_	+
3	T2CNoMo	PLND 5/93	4.5	0.1	· •	+
4	T2BNoMo	RRP 3/92	NMA	0.4	_	+
5	T3NxMo	Proscar + Flutamide	51.3	1.0	-	+
6	Recur T3	I-125 1986	54.7	1.4	-	+
7	T3ANoMo	RRP 10/92	NMA	0.3	_	+
8	ТЗМхМо	XRT 1987	7.5	0.1	-	<b>-</b> ·
9	T3NxMo	Proscar + Flutamide	35.4	0.7	_	-
10	D2	S/P XRT Flutamide +Emcyt	311	4.5	+	+
11	D2	RRP 4/91 Lupron 10/92 Velban + Emcyt 12/92	1534	1.4	+	+
12	T2NoMo	RRP 8/91	NMA	0.5	-	.+
13	ТЗМоМо	RRP 1/88 Lupron + Flutamide 5/92	0.1	0.3	<del>-</del>	~
14	D1	PLND 1989 XRT 1989	1.6	0.4	-	-
15	D1	Proscar + Flutamide	20.8	0.5	<del>-</del>	-
16	T2CNoMo	RRP 4/92	0.1	0.3	-	-

#### FIGURE 31A

	10	20	30	40	50	. 60
1	AAGGGTGCTC	CTTAGGCTGA	ATGCTTGCAG	ACAGGATGCT	TGGTTACAGA	TGGGCTGTGA
	TTCCCACGAC	GAATCCGACT	TACGAACGTC	TGTCCTACGA	ACCAATGTCT	ACCCGACACT
61	CTCGAGTGGA	GTTTTATAAG	GGTGCTCCTT	AGGCTGAATG	CTTGCAGACA	GGATGCTTGG
	GAGCTCACCT	CAAAATATTC	CCACGAGGAA	TCCGACTTAC	GAACGTCTGT	CCTACGAACC
121	TTACAGATGG	GCTGTGAGCT	GGGTGCTTGT	AAGAGGATGC	TTGGGTGCTA	AGTGAGCCAT
	AATGTCTACC	CGACACTCGA	CCCACGAACA	TTCTCCTACG	AACCCACGAT	TCACTCGGTA
181	TTGCAGTTGA	CCCTATTCTT	GGAACATTCA	TTCCCCTCTA	CCCCTGTTTC	TGTTCCTGCC
	AACGTCAACT	GGGATAAGAA	CCTTGTAAGT	AAGGGGAGAT	GGGGACAAAG	ACAAGGACGG
241	AGCTAAGCCC	ATTTTTCATT	TTTCTTTTAA	CTCCTTAGCG	CTCCGCAAAA	CTTAATCAAT
	TCGATTCGGG	TAAAAAGTAA	AAAGAAAATT	GAGGAATCGC	GAGGCGTTTT	GAATTAGTTA
301	TTCTTTAAAC	CTCAGTTTC GAGTCAAAAS	TTATCTGTAA AATAGACATT	AAGGTAAATA TTCCATTTAT	ATAATACAGG TATTATGTCC	GTGCAACAGA CACGTTGTCT
361	AAAATCTAGT	GTGGTTTACA	TAATCA CCTG	TTAGAGATTT	TAAATTATTT	CAGGATAAGT .
	TTTTAGATCA	CACCAAATGT	ATTAGTGGAC	AATCTCTAAA	ATTTAATAAA	GTCCTATTCA
	GTACTATTAA	AAATGAAATA TITACTTTAT	TACGTGTATT	TCGTGTATCA	CACCACAGGA	GGTATATCTT
	TTACGAGTCA	ATATTGGTTA TATAACCAAT	AATTGATGAA	CAACTTCCAA	ATAGAAGAGG	TGATTTGACA
541	AAGTTCCACA	AGCCTTACAA	TATGTGACAG	ATATTCATTC	ATTGTCTGAA	TTCTTCAAAT
	TTCAAGGTGT	TCGGAATGTT	ATACACTGTC	TATAAGTAAG	TAACAGACTT	AAGAAGTTTA
	TGTAGGAGAA	CACCATAGCG GTGGTATCGC	AGAATAATTA	ACTTAATAAT	TAACTTATTT	AAGATAACAA
	GTTTTTAGTG	TTTTATATTT	TIGACTITAA	ACGAATGAAT	ATTAGTGTAG	ATTGGAAGTT
721	AGAAAACACA	TTAACCAACT	GTACTGGGTA	ATGTTACTGG	GTGATCCCAC	GTTTTACAAA
	TCTTTTGTGT	AATTGGTTGA	CATGACCCAT	TACAATGACC	CACTAGGGTG	CAAAATGTTT

#### FIGURE 31B

781	TGAGAAGATA	A TATTCTGGTA	AGTTGAATAC	TTAGCACCCA	GGGGTAATCA	GCTTGGACAG
	ACTCTTCTAT	I ATAAGACCAT	TCAACTTATG	AATCGTGGGT	CCCCATTAGT	CGAACCTGTC
841	GACCAGGTCC	AAAGACTGTT	AAGAGTCTTC	TGACTCCAAA	CTCAGTGCTC	CCTCCAGTGC
	CTGGTCCAGC	TTTCTGACAA	TTCTCAGAAG	ACTGAGGTTT	GAGTCACGAG	GGAGGTCACG
901	CACAAGCAAI	CTCCATAAAG	GTATCCTGTG	CTGAATAGAG	ACTGTAGAGT	GGTACAAAGT
	GTGTTCGTTT	GAGGTATTTC	CATAGGACAC	GACTTATCTC	TGACATCTCA	CCATGTTTCA
961	AAGACAGACA	TTATATTANG	TCTTAGCTTT	GTGACTTCGA	ATGACTTACC	TAATCTAGCT
	TTCTGTCTGT	AATATAATTC	AGAATCGAAA	CACTGAAGCT	TACTGAATGG	ATTAGATCGA
1021	AAATTTCAGT	TTTACCATGT	GTAAATCAGG	AAGAGTAATA	GAACAAACCT	TGAAGGGTCC
	TTTAAAGTCA	AAATGGTACA	CATTTAGTCC	TTCTCATTAT	CTTGTTTGGA	ACTTCCCAGG
1081	CAATGGTGAT	TAAATGAGGT	GATGTACATA	ACATGCATCA	CTCATAATAA	GTGCTCTTTA
	GTTACCACTA	ATTTACTCCA	CTACATGTAT	TGTACGTAGT	GAGTATTATT	CACGAGAAAT
1141	AATATTAGTC	ACTATTATTA	GCCATCTCTG	ATTAGATTTG	ACAATAGGAA	CATTAGGAAA
	TTATAATCAG	TGATAATAAT	CGGTAGAGAC	TAATCTAAAC	TGTTATCCTT	GTAATCCTTT
1201	GATATAGTAC	ATTCAGGATT	TTGTTAGAAA	GAGATGAAGA	AATTCCCTTC	CTTCCTGCCC
	CTATATCATG	TAAGTCCTAA	AACAATCTTT	CTCTACTTCT	TTAAGGGAAG	GAAGGACGGG
1261	TAGGTCATCT	AGGAGTTGTC	ATGGTTCATT	GTTGACAAAT	TAATTTTCCC	AAATTTTTCA
	ATCCAGTAGA	TCCTCAACAG	TACCAAGTAA	CAACTGTTTA	ATTAAAAGGG	TTTAAAAAGT
1321	CTTTGCTCAG	AAAGTCTACA	TCGAAGCACC	CAAGACTGTA	CAATCTAGTC	CATCTTTTTC
	GAAACGAGTC	TTTCAGATGT	AGCTTCGTGG	GTTCTGACAT	GTTAGATCAG	GTAGAAAAAG
	GTGAATTGAG	ATACTGTGCT TATGACACGA	GAGGGAAAGA	GTTTCGTTTG	ACAAACGATA	AGGAACTTAT
	GTGAGACTCA	TTTCTGCCTT AAAGACGGAA	ACGGATGAGT	CGACCGGGTA	CCGGGGATTA	CAAAGAAGAG
	TAGAGGTGAC	GGTCAAATCC CCAGTTTAGG	ATGGACATGG	AATACCAAGA	CAATTTTCGT	CACGAAGGTA
1561	AAAGTACTCC	TAGCAAATGC	ACGCCCTCTC	TCACGGATTA	TANGANCACA	GTTTATTTTA

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#### FIGURE 31C

TTTCATGAGG ATCGTTTACG TGCCGGAGAG AGTGCCTAAT ATTCTTGTGT CAAATAAAAT 1621 TAAAGCATGT AGCTATTCTC TCCCTCGAAA TACGATTATT ATTATTAAGA ATTTATAGCA ATTTCGTACA TCGATAAGAG AGGGAGCTTT ATGCTAATAA TAATAATTCT TAAATATCGT 1681 GGGATATAAT TTTGTATGAT GATTCTTCTG GTTAATCCAA CCAAGATTGA TTTTATATCT CCCTATATTA AAACATACTA CTAAGAAGAC CAATTAGGTT GGTTCTAACT AAAATATAGA 1741 ATTACGTAAG ACAGTAGCCA GACATAGCCG GGATATGAAA ATAAAGTCTC TGCCTTCAAC TAATGCATTC TGTCATCGGT CTGTATCGGC CCTATACTTT TATTTCAGAG ACGGAAGTTG TTCAAGGTCA TAAGAAAAGA AAGGAGGGGA GGGAAGGGGA GGGGAAGGAA 1861 CCCTTTCCCT TCCCTTCCTT TCTTTCTTGA GGGAGTCTCA CTCTGTCACC AGGCTCCAGT GGGAAAGGGA AGGGAAGGAA AGAAAGAACT CCCTCAGAGT GAGACAGTGG TCCGAGGTCA 1921 GCAGTGGCGC TATCTTGGCT GACTGCAACC TCCGCCTCCC CGGTTCAAGC GATTCTCCTG CGTCACCGCG ATAGAACCGA CTGACGTTGG AGGCGGAGGG GCCAAGTTCG CTAAGAGGAC 1981 CCTCAGCCTC CTGAGTAGCT GGGACTACAG GAGCCCGCCA CCACGCCCAG CTAATTTTTG GGAGTCGGAG GACTCATCGA CCCTGATGTC CTCGGGCGGT GGTGCGGGTC GATTAAAAAC 2041 TATTTTAGT AGAGATGGGG TTTCACCATG TTGGCCAGGA TGGTCTCGAT TTCTCGACTT ATAAAAATCA TCTCTACCCC AAAGTGGTAC AACCGGTCCT ACCAGAGCTA AAGAGCTGAA 2101 CGTGATCCGC CTGTCTGGGC CTCCCAAAGT GCTGGGATTA CAGGCGTGAG CCACCACGCC GCACTAGGCG GACAGACCCG GAGGGTTTCA CGACCCTAAT GTCCGCACTC GGTGGTGCGG 2161 CGGCTTTAAA AAATGGTTTT GTAATGTAAG TGGAGGATAA TACCCTACAT GTTTATTAAT GCCGAAATTT TTTACCAAAA CATTACATTC ACCTCCTATT ATGGGATGTA CAAATAATTA 2221 AACAATAATA TTCTTTAGGA AAAAGGGCGC GGTGGTGATT TACACTGATG ACAAGCATTC TTGTTATTAT AAGAAATCCT TTTTCCCGCG CCACCACTAA ATGTGACTAC TGTTCGTAAG 2281 CCGACTATGG AAAAAAAGCG CAGCTTTTTC TGCTCTGCTT TTATTCAGTA GAGTATTGTA GGCTGATACC TTTTTTTCGC GTCGAAAAAG ACGAGACGAA AATAAGTCAT CTCATAACAT 2341 GAGATTGTAT AGAATTTCAG AGTTGAATAA AAGTTCCTCA TAATTATAGG AGTGGAGAGA CTCTAACATA TCTTAAAGTC TCAACTTATT TTCAAGGAGT ATTAATATCC TCACCTCTCT

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#### FIGURE 31D

2401		TTTCTTCCTT		
2461		TTTTTAAGGC AAAAATTCCG		
2521		TCTCTCTCGC. AGAGAGAGCG		
2581		ACCCCAGGTC TGGGGTCCAG		
2641		GAGAGAGACT CTCTCTCTGA		
2701		GCGGGTCCCG CGCCCAGGGC		
2761		ACTCGGCTGT TGAGCCGACA		
2821		CGGGTGGCTT GCCCACCGAA		
2881		TCGGAGTCTT AGCCTCAGAA		
2941		CCTGTTGCTG GGACAACGAC		
3001	GGTGAGCACC			

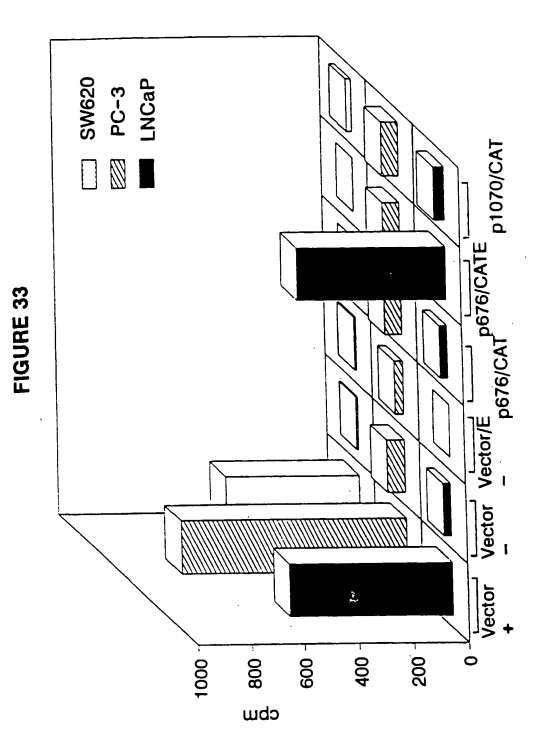
#### FIGURE 32

#### Potential binding sites on the PSM promoter\*

Site	Seq	**Location	#nt matched	
AP1	TKAGTCA	1145	7/7	
E2-RS	ACCNNNNNNGGT	1940 1951	12/12 12/12	
GHF ·	NNNTAAATNNN	580 753 1340 1882 1930 1979 2001 2334 2374 2591 2620 2686	11/11 11/11 11/11 11/11 11/11 11/11 11/11 11/11 11/11 11/11	
JVC repeat	GGGNGGRR	1165 1175 1180 1185 1190	8/8 8/8 8/8 8/8 8/8	
NFkB	GGGRHTYYHC	964	10/10	
uteroglobi	RYYWSGTG	250 921 1104	8/8 8/8 8/8	
IFN AAW	AANGAAAGGR590	13/13	Cell 41:509 (1985)	

<sup>\*</sup> the PSM promoter sequence 683XFRVS (Fig. 1) starts from the 5' end of the promoter fragment. The 3' region overlapps the previously published PSM cDNA at nt#2485,i.e. the putatative transcription start site is at nt#2485 on sequence 683XFRVS. \*The number refered to in this table is in reference to sequence 683XF107 which is the complement and inverse of 683XFRVS.

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COCOOTIGET TO AGAGGGCGCCAGT AGAGCAGCAGCAGGCGCGGGTCCCGGGAGGCCGGCTCTGCTCGCGCCGAG OBACCCC ADD TCTUBADCOAATTCCA UCCTBCAUDDCTDAIAADCOADDCATTAUTBAUATTBADADACTTTACCC TGT TUBARUCABATOTTOCCTCTCTCTCUCTCUUATTOOTTCABTOCACTCTABAAAAACTGCTGTOOTGGAAAAACT

CTCAAAAGGGGCCGGATTTCCT

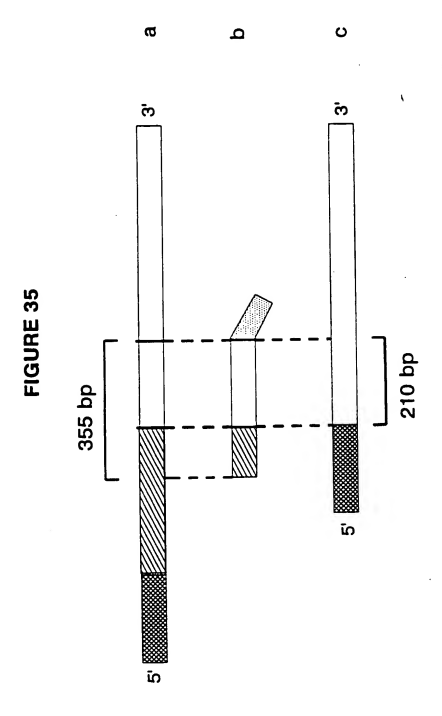
FIGURE 34

ATO TOO AAT CTC CTT CAC DAA ACC DAC TCO DCT OTO DCC ACC DCO COC COC CCO COC TOO CTO Pro Arg Trp Leu Arg Arg **-**-Vel Ale Ser Ale Asp Glu Thr Met Trp Aen Leu Leu Hie TOC OCT DOG OCO CTO OTO CTO OCO OOT OUCTIC III CTC CTC GOC TTC CTC TTC OOA TOO III Gly Phe Phe Leu Leu Gly Phe Leu Phe Gly Leu Vai Leu Ain Gly ij al, Cye Ale

ATA AAA TOC TOC AAT BAA BOT ACT AAC ATT ACT COA AAB CAT AAT ATB AAA BOA TIT TIB BAT BAA 3 Als Thr Asn 11e Thr Pro Lys 11is Asn Met Lys Als Phe Leu Asp Ser Aen Glu 110 Lys Ser

Ī TOO AAA OCT OAO AAC ATC AAO AAU TTC TTA TAT AAT TTT ACA CAU ATA CCA CAT TTA GCA GOA c C Ale Glu Aen 116 Lys Lys Phe Leu Tyr Asn Phe Thr Gln 116 Pro 1118 Leu Ais

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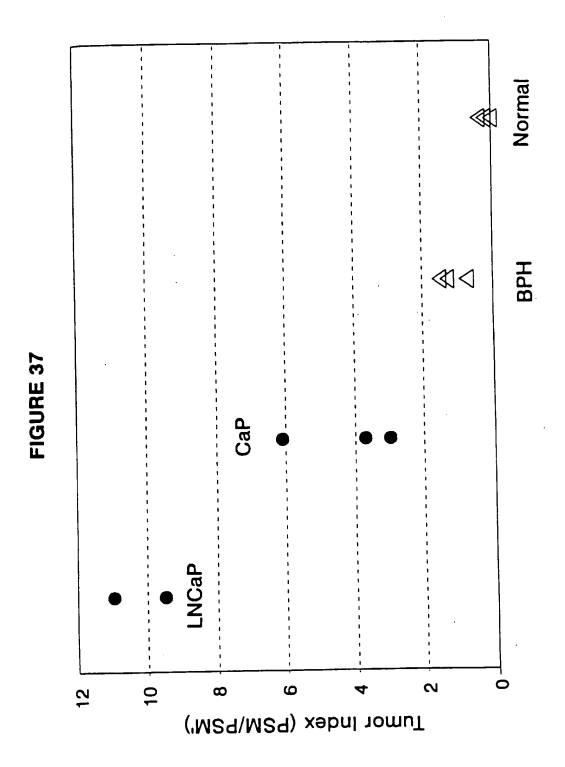


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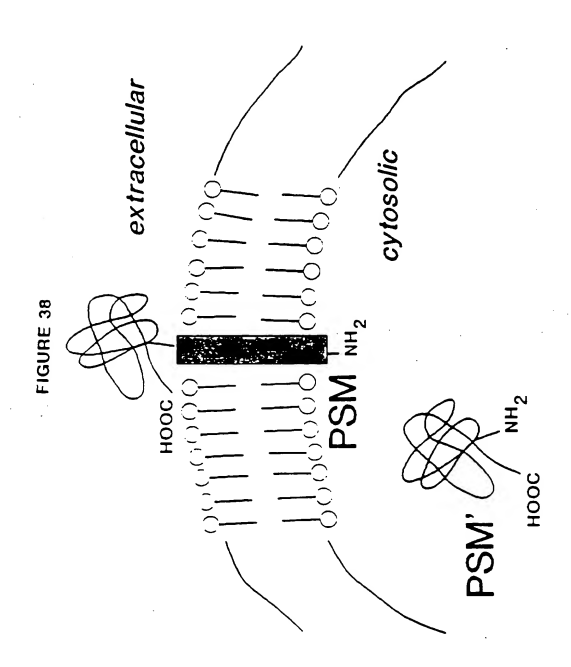


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#### FIGURE 39

	1	l 20	30	40	50	60
1	TTTGCAGAC	TGACCAACTI	TCTAAGAAAA	GCAGAACCAC	ACAGGCAAGC	TCAGACTCTT
		A ACTGGTTGAA				
61	TTATTAAAT AATAATTTA	CCAGTTTTGA GGTCAAAACT	CTTTGCCACT	TCTTAGTGGC	CITGAACAAG	TTACCGAGTC
121			•			
121	GAGAGTCGCA	TAGTTACCCT ATCAATGGGA	TAAAATTACT	ACTCCTATTA	TAATAGACGG	CAAATTATTG
181	GTATAGTAAA	TATATAGCAT	GTAAATCTCC	TAGCAGAGTA	CTGGGATTTC	GCCACTTTAT
	CATATCAPPI	ATATATCGTA	CATTTAGAGG	ATCGTCTCAT	GACCCTAAAG	CGGTGAAATA
241	TTCTTCTTTA	CCAAGATACT	CCTATTGGAC	TTAATACACA	GGACTAGTCT	AAGGTATCAC
		GGTTCTATGA		•	-	
301	CAGGTAGTCC GTCCATCAGG	ACTOCTGCTC TGAGGACGAG	GGAATCTGAC	CCGGGATTAG	AGTAGGGCAT	GGACCAGATG
	•	•				
361	GGTTTAAACA CCAAATTTGT	AATTCAATAT TTAASTTATA	CTTCCACTAG	CTTCACCTTG	GGGTTGTAAA	AGTTTTTGAA
421	DIACACACTG DETETETGAC	TGCTCATAAC ACGAGTATTG	AATCTTCATC	TCTTAAAAGG	ATTITATION	TCCTGGTATC
			•			
481	GAGTGAGAGT	TCCCTTGTAT AGGGAACATA	TCCGTGCTCA AGGCACGAGT	GTGGCTGACA	CAGAAGAGTT	CTTTATHNHH
541	HARRESTER HER	CATCCTGTTC GTAGGACAAG	ATTTTTCAGA	TCTCAGTTCA	AGCATCTCGT	CCTCAGTGTG
					•	
601	GTGTTNNCTG	ATCCCTCACT TAGGGAGTGA	CTAATCCAAG	TCTTTCTGTT	TTATGCACAG	GTTGGAATCT
661	TATTTCCGTT	TGCGNNCCAA ACGCNNGGTT	TCHAATHGTA	TTTAATATGC	ATGTATATAT	GTATGTGCAT
721	TTGTATGCTA AACATACGAT	NGCGATTAAG NCGCTAATTC	AACTAGAATA TTGATCTTAT	ATTAATAATT TAATTATTAA	GGAAGTCTAG	AAGTGG

#### FIGURE 40A

	1	0 2	0 30	4.0	50	60
	1 TGAAAAATA	C ATCAAAAAT.	i A ggcatgagat	   ACGAGCCTAT	AGATAGGACT	:
	ACTTTTAT	G TAGTTTTA	T CCGTACTCTA	TGCTCGGATA	TCTATCCTGA	ATAAAAATA
6	TATTGTTGT.	A TGTATTATT	T GTAAAACACA	AATTATCAAT	ATTACCTCTG	ACATTAGGTG
	ATAACAACA'	I ACATAATAA	A CATTTTGTGT	TTAATAGTTA	TAATGGAGAC	TGTAATCCAC
123	AGATATTCT	AATTTTAAT	Tererrecer	ACTITCACTO	AAAAAGAGTO	ATCCASSOS
	TCTATAAGA	TTAAAATTAJ	A AGAGAACGGA	TGAAAGTGAC	TTTTTCTCAG	TACGTTTGTC
181	ATTTTTAAGT	TGCAAACCAA	TTGCAAAATA	TTTTTTTATC	CAACTTCALT	GATA CCTA TO
	TAAAAATTC	ACGTTTGGTT	AACGTTTTAT	AAAAAAATAG	GTTGAAGTTA	CTATCCATAA
241	GCTGTTAATT	CTAAGATATO	CATTAATTGT	TTCAACTAAT	GGGTGTCAAA	CGAGATCTTC
	CGACAATTAA	GATTCTATAC	GTAATTAACA	AAGTTGATTA	CCCACAGTTT	GCTCTACAAG
301	TGAAAATGAA	GGCAAAAAGG	AGATOCACCT	TOTACTTTCA	TAAAGT	A T
	ACTITIACTI	CCGTTTTTCC	TCTAGGTGGA	AGATGAAAGT	ATTTCAAAGA	TAGAAGGAGA
361	GCTGACTCAA	ATAAGCATTT	AATACATTTT	ATAACGAATT	AATTATGAAT	A T 2
	CGACTGAGTT	TATTOSTAAA	TTATGTAAAA	TATTGCTTAA	TTAATACTTA	TATAAAGTTT
421	TAAATAAATT	ATTTCCAAGT	GTTGAAGGAA	ATTCASACTT	CT-3 3 TYPE-C-	CTC) TECTO
	ATTTATTTAA	TAAAGGTTCA	CAACTTCCTT	TAAGTCTGAA	GATTAAACGA	GACTAAGACT
481	AACTAAAACA TTGATTTTGT	AATGCTCTGT	GAGAGTTTGC	GTTTCCAGTG	1157150075	161112000
	TTGATTTTGT	TTACGAGACA	CTCTCAAACS	CAAAGGTCAC	TTCATCGCAC	TCTTTAGGTT
541	GTCAGACAGC	TACATGAAAC	TACATTTATT	AGCTCTCTGC	CFTFCVCCVC	7551001710
	CAGTCTGTCG	ATGTACTITG	ATGTAAATGG	TCGAGAGACG	GTCTGTGGTC	ACGTGCTATC
601	CGCAGAACAT	GTAGCTAGAT	CTCACTCATA	COTIVINADA		
	GCGTCTTGTA	CATCGATCTA	GAGTCAGTAT	ССЕМИНИМИМ	ининининин	AGACCTTGCA TCTGGAACGT
661	GTTGGCTTTT	AACCTGAAGG	AGATARCCC	1617700100	Chara Land	
	CAACCGAAAA	TTGGACTTCC	TCTATTCCGT	TCTAAGGTCC	CAAATAAATC	AGAAATTACA TCTTTAATGT
721	GGATCTGGGA	ATAAAGTAGT	TACAAAATTA ATGTTTTAAT	GTCCCCAACC	AGCTTTCATG	GAGCTTTCAA
		-UTITIONICA	AIGITITAAT	CAGGGGTTGG	TCGAAACTAC	CTCC 3 3 3 COM

#### FIGURE 40B

			1 WY 1 COCH 1 C	CATACAATGC	ACATACATAT	ATACATGC AT
	TAATTAATAA	AAGATCAAGA	ATTAGCGTAC	GTATGTTACG	TGTATGTATA	TATGTACGTA
841	ATTAAAATAC	ATGATTGGAC	CCAAACCCAA	1711517700	1000000000	
• • •	TAATTTATO	TACTAACCTG	CCTTTCCCTT	MIAAGATICC	ACCIGIGCAT	AAAACAGAAA
	179.1	· inclinacio	COTTIGCCTT	TATTCTAAGG	TGGACACGTA	TITTGTCTT:
901	GACTTGGTTA	GAGTGAGGGA	TCAGGAAACA	CCACACTGAG	GACGAGATON	NENNYMMAN
	CTGAACCAAT	CTCACTCCCT	AGTECTTTGT	GGTGTGACTC	CTGCTCTACN	инининини
				•		
961	NTAGTGGGTG	GGGGGGGGAC	ATCAATAAAG	AACTCTTCTG	TGTCAGCCAC	TGAGCACGGA
	NATCACCCAC	cccccccrc	TAGTTATTTC	TTGAGAAGAC	ACASTCGGTG	ACTOSTGOOT
1021	ATAAAGGGAT	GAGASTGASG	GCAANTACCA	GAAGAATAAA	AFCCTTTTAA	GAGATCAACA
	TATTTCCCTA	CTCTCACTCC	CGTTNATGGT	CITCITAITI	TAGGAAAATT	CTOTACTTO
	٠.					
1081	TIGTTATGAG	CACAGTGTGT	GGHTTCAAAA	ATCTTTTAAC	AACCCCAAGG	TGAAGCTACT
	AACAATACTC	STSTCACACA	CCNAAGTTTT	TAGAAAATTG	TTGGGGTTCC	ACTTCGATCA
1141	TGGAAGATAT	TTSAATTTGT	TTAAACCCAT	CTGGTCCTAG	CCCTATTCTT	TGAATCCGA
	ACCTTCTATA	AACTTAAACA	AATTTGGGTA	GACCAGGATC	GGGATAAGAA	ACTTAGGCTT
1201	GAGGTCAAGA	ATTOCGASCA	GASTSSACTA	CCTGTGATAC	CTTAGACTAG	TOTETETAT
	CTCCAGTTCT	TAAGGCTCGT	CTCACCTGAT	GGACACTATG	GAATCTGATC	AGGACACATA
1261	TCAAGTCCAA	TGAGAGTATC	TGTAAGAGAA	TAAGTGCGAA	ATCCAGATCT	
	AGTTCAGGTT	ACTOTOATAG	ACATTETETT	ATTCACGCTT	TAGGTCTAGA	

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#### FIGURE 41

	10	ī	30	40	50 i	<b>6</b> 0
1	GGATTCTGTT	GAGCCCTAGC	TCATTATGAT	GTCCTGTTGT	CCTACCCAAA	TAAGACTCAT
	CCTAAGACAA	CTCGGGATCG	AGTAATACTA	CAGGACAACA	GGATGGGTTT	ATTCTGAGTA
61	CCCAACTACA	TCTCAATAAT	TAATGAAGAT	GGAAATGAGG	TAAAAAATAA	TAAATAAAT
	GGGTTGATGT	AGAGTTATTA	ATTACTTCTA	CCTTTACTCC	ATTTTTTTATT	ATTTATTTA
121	AAAAGAAACA	TTCCCCCCA	TTTATTATTT	TTTCAAATAC	CTTCTATGAA	ATAATGTTCT
	TTTTCTTTGT	AAGGGGGGGT	AAATAATAAA	AAAGTTTATG	GAAGATACTT	TATTACAAGA
181	ATCCCTCTCT	TAATATAAA	AGAAATCAAT	ATTATTGGAA	CTGTGAATAC	CTTTAATATC
	TAGGGAGAGA	ATTAAATTA	TCTTTAGTTA	TAATAACCTT	GACACTTATG	GAAATTATAG
241	TCATTATCCG	GTGTCAACTA	CTTTCCTATG	ATGTTGAGTT	ACTGGGTTTA	GAAGTCGGGA
	ASTAATAGGC	CACAGTTGAT	GAAAGGATAC	TACAACTCAA	TGACCCAAAT	CTTCAGCCCT
301	AATAATGCTG	TAAANNANAN	AGTTAGTCTA	CACACCAATA	TCAAATATGA	TATACTTGTA
	TTATTATGAC	ATTTA	TCAATCAGAT	GTGTGGTTAT	AGTTTATACT	ATATGAACAT
361	AACCTCCAAG	CATAAAAAGA	GATACTTTAT	AAAAGAGGTT	CTTTTTTTCT	TTTTTTTTTT
	TTGGAGGTTC	GTATTTTTCT	CTATGAAATA	TTTTCTCCAA	GAAAAAAAGA	AAAAAAAA
421	TODAGATGGA	GTTT DACTOC	TGTCAGGCAG	GENGAGTGCA	GTGGTGCCAT	CTCGGCTCAC
	AGGTCTACCT	CAAAGTGAGG	ACAGTCCGTC	CGNCTCACGT	CACCACGGTA	GAGCCGAGTG
461	TGCAACCTCC	ACCTCCCATG	TT LAAJGGAT	TCTCCTTCCT	CAGTCTCCTG	AGTAGCTGGG
	ACGTTGGAGG	TGGAGGGTAC	AAGTTCCCTA	AGAGGAAGGA	GTCAGAGGAC	TCATCGACCC
541	ATTACAGGTG	TGCACCACCA	CACCCASCTA	ATTTTTGTAT	TTTTAATAGA	GACAGGGTTT
	TAATGTCCAC	ACGTGGTGGT	STSGGTCGAT	TAAAAACATA	AAAATTATCT	CTGTCCCAAA
601	CSATCGATGT	TGGCCAGGCT	AGTCTCGAAC	TCCTGACCTC	TAGGTGATCC	ACCCGCTCAG
	GCTASCTAIA	ACCGGTCCGA	TCAGAGCTTG	AGGACTGGAG	ATCCACTAGG	TGGGCGAGTC
661	CTCCCAAAGT	TGTAGAATTA	CACGTGTGAG	GCACTGCGCC	TTGCCAGGAG	ATACATTTT
	GAGGGTTTCA	ACATCTTAAT	GTGCACACTC	CGTGACGCGG	AACGGTCCTC	TATGTAAAA
721	GATAGGTTTA	ATTTATAAAG	ACACTGCACA	GATTTGAGTT	GCTGGGAAAT	GCACGGATTC
	CTATCCAAAT	TAAATATTTC	TGTGACGTGT	CTAAACTCAA	CGACCCTTTA	CGTGCCTAAG

781 CAGTATGCA GTCATACGT

AAACAGTTAA AGTTTAATTA CTATAATGAA ACACAAAAAA AATGAATATT TTAGTTTTAT 1"FIGTGAATT TCAAAGTAAT GATATTAGTT TGTGTTTTT TTACTTAAA 50 0 30 AATCAAAATA

FIGURE 42

GTATCAGATA TCAGTAGAGG CTCAATCAAT CCTTCAGGAT TTTGATGATA TAGAAAATAC AGTCATCTCC CACTTACTTA GGAACTCCTA AAACTACTAT ATCTTTTATG 9

CATAGTCTAT

CCCAGCACTA TGCTAGAAGT TGTGAAGAAT TGAGGAGATG AATAAATCAC AGATTCTGTC GGGTCGTGAT ACGATGTTCA AGACTTGTTA AGTGCTGTAC TTATTTAGTG TGTAAGACAG 121

TTAGATCTAT TCAGGAAACA AAGCTAAAAA AACCCCACCA ATAACTAAAA TATTGATTTT GAGTITTACC AATCTAGATA AGTCCTTTTGT TTCGATTTTT TTGGGGTGGT CTCAAAATGG 181

TCTTTTCGAG AGANAAGCTC ATCAACCAAA TGAAAAACAA CAATCATAAA ATAAGTAAGT ACCTATAGAA TAGTTGGTTT ACTTTTTGTT GTTAGTATT TATTCATTCA TGGATATCTT 241

TITTCTTAGA GGAATTTTCC TTATGATATA TGACATTTTG ACACTGACTA TGTGACTGAT 301 AGAGGAGGTA AAAAGAATCT CCTTAAAAGG AATACTATAT ACTGTAAAAC TCTCCTCCAT

AGAAGGNA TCTTCCTT 190

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#### FIGURE 43A

	10	`2¢	30	40	50	60
1	TATGGGAAAS	TTTTCAGAGG	AAATAAGGTA	AGGGAAAAGT	TATCTCTTTT	TTTCTCTCCC
	ATACCCTTTC	AAAAGTCTCC	TTTATTCCAT	TCCCTTTTCA	ATAGAGAAAA	AAAGAGAGGG
61	CCAATGTAAA	AAGTTATAGT	GGGTTTTACA	TGTGTAGAAT	CATTTTCTTA	AAACTITATG
	GGTTACATTT	TTCAATATCA	CCCAAAATGT	ACACATCTTA	GTAAAAGAAT	TTTGAAATAC
121	AATACCATTA	TTTTCTTGTA	TTCTGTGACA	TGCCACCTTA	CAGAGAGGAC	ACATTTACTA
	TTATGGTAAT	AAAAGAACAT	AAGACACTGT	ACGGTGGAAT	GTCTCTCCTG	TGTAAATGAT
181	GGTTATATCC	CGGGGTTAAA	TTCGAGCATT	GGAATTTGGC	CAGTGTAGAT	GTTTAGAGTG
	CCAATATAGG	GCCCCAATTT	AAGCTCGTAA	CCTTAAACCG	GTCACATCTA	CAAATCTCAC
241	AACAGAACAA	TTTTTCTGTG	CTTACAGGTT	ATGGCTGTGG	CGTA IAAGAA	GCATGCACTG
	TIGICTIGIT	AAAAAGACAC	GAATGTCCAA	TACCGACACC	GCATGTTCTT	CGTACGTGAC
301	GGTTTATTAT	TAACTTTCAG	TATCTTTGTT	TTAAATATTT	TOTACAAAA	TGTTTACTAA
	CCAAATAATA	ATTGAAAGTC	ATAGAAACAA	AATTTATAAA	AGATGTTTTT	ACAAATGATT
361	ATTAAATTGT TAATTTAACA	AGTATGAATT TCATACTTAA				
421	AAAAATTACT	GTCATTTGAT	TTGTTAATAT	ATTTTTCTCT	TTAGTGGGAA	ATTAAATTAA
	TTTTTAATGA	CAGTAAACTA	AATAATTATA	TAXAAAGAGA	AATCACCCTT	TAATTTAATT
461	AAAATTCCTT	TOGACTOTCA	GACAATAGGA	TIGCTGTGGT	CTACTTGCTT	ATTATATTG
	TTTTAAGGAA	AGCTGACAGT	CTGTTATCCT	AACGACACCA	GATGAACGAA	TAATATAAAC
541	TAGAGTCTAG ATCTCAGATC	AATGCAATCT TTACGTTAGA				
601	TGAGAAACTA	TTCCAGACCT	COTTATOSGO	TTAGCCAAGG	NTATCCTTCA	GCTGGCATTG
	ACTCTTTGAT	AAGGTCTGGA	GGAATACCCS	AATCGGTTCC	NATAGGAAGT	CGACCGTAAC
661	CAGGGTGACT	TCTNCCTCHN	AATCCAGCTC	TCTNTCACAG	ATGTGATCCA	AGAGACACTC
	GTCCCACTGA	AGANGGAGHN	TTAGGTCGAG	AGANAGTGTC	TACACTAGGT	TCTCTGTGAG
721	ACAATTAATC TGTTAATTAG					

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#### FIGURE 43B

- TSI TGAAGCTTIN NTCACTGTCA ATTCTGATCA GATATATGAC AATTTTAAAT TATTTGCAGT ACTTCGAAAN NASTGACAGT TAAGACTAGT CTATATACTG TTAAAATTTA ATAAACGTCA
- 841 GTGTAAGAAA CGCTTCAGGT AGTTTAAATT TAAGGCT CACATTCTTT GCGAAGTCCA TCAAATTTAA ATTCCGA

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#### FIGURE 44A

	10	20	30	40	50	60
1	CTCCTTTGGC	CCCTGCCAGC	TGGGCATTTT	TAACCTAGTT	TACACAGTGT	CTTTTTTTCC
	GAGGAAACCG	GGGACGGTCG	ACCCGTAAAA	ATTGGATCAA	ATGTGTCACA	GAAAAAAAGG
€1	AAATTTAAA	TTGGTTGTTC	CAGATTCGGT	AATATCAATT	TTTATATTA	CACTTAAATG
	TTTAAAATAA	AACCAACAAG	GTCTAAGCCA	TTATAGTTAA	AAATTATAAA	GTGAATTTAC
121	AGTACCAGAA	CTTTATCTTC	AACCTTTTTC	TCATTAGGCC	TACAACATAG	GACATCTCGG
	TCATGGTCTT	GAAATAGAAG	TTGGAAAAAG	AGTAATCCGG	ATGTTGTATC	CTGTAGAGCC
181	ATAGAATTTC	CTTTTCTTTT	TGCTACTATA	AGCTGCTAAA	ATCCTCAGAA	CATCAGATTT
	TATC <b>TTAAA</b> G	GAAAAGAAAA	ACGATGATAT	TCGACGATTT	TAGGAGTCTT	GTAGTCTAAA
241	AGAAATGTTC	TTATTAGTGG	TAGTGAGCAT	TTGCTATTTC	CTACCACTAG	CTTACAAATA
	TCTTTACAAG	AATAATCACC	ATCACTCGTA	AACGATAAAG	GATGGTGATC	GAATGTTTAT
391	TAATAAGCAA	GTAGACCCCA	CAGGOCAAAT	TCCTATTTGT	TCTACAGTCG	AAAGGGAATT
	ATTATTCGTT	CATCTGGGGT	GTCCGGTTTA	AGGATAAACA	AGATGTCAGC	TTTCCCTTAA
361	TTTTAAAATT	TAATTTCCAC ATTAAAJGTG	TAAAGAGAAA ATTTCTCTTT	AATATATTAA TTATATAATT	CAATCAAATT GTTAGTTTAA	GACAGTCGAT CTGTCAGCTA
421	TTTAATTSCT	ATGTGTAATT	GTTTTCCCTC	ATTATTTATA	ACAATTCATA	CTACAATTTA
	AAATTAACGA	TAGAGATTAA	TAAAAGSSAS	TAATAAATAT	TGTTAAGTAT	GATGTTAAAT
451	ATTTASTARA	CATTTTTGTA	GACCATATTT	AAAACAAAGA	TACTGAAAGT	TAATATAAAC
	TARATCATTT	GTAAAAACAT	CTGGTATAAA	TTTTGTTTCT	ATGACTTTCA	ATTATATTTG
[4]	CCAGTGCATG	CTCTCTGTAG	GECACAGCCA	TAACCTGTAA	GCACAGAAAA	ATTTGTTCTG
	GGTCACGTAC	GAGAGACATO	CGSTSTCGGT	ATTGGACATT	CGTGTCTTTT	TAAACAAGAC
601	TTACTCTAAA	CATCTAIACT	GGCCAAATTC	CAATGCTCGA	ATTTAACCCC	GGGATATAAC
	AATGAGATTT	GTAGATITGA	CCGGTTTAAG	GTTACGAGCT	TAAATTGGGG	CCCTATATTG
661	CTASTAAATG	TGTCCTCTCT	GTCAASGTGG	GCATGTCACA	GAATACAGAA	CAATCAATGG
	SATCATTTAC	ACAGGAGAGA	CAGTTCCACC	CGTACAGTGT	CTTATGTCTT	GTTAGTTACC
721	TATTCATAAA	GTTTTAAGAA	AATGATTCTA	CACATGTAAA	ACCCACTATA	ACTTTTTACA
	ATAAGTATTT	CAAAATTCTT	TTACTAAGAT	GTGTACATTT	TGGGTGATAT	TGAAAAATGT

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#### FIGURE 44B

E41 CATATCTOOC AATTACAATT TTOOCAGASC AATTGATTTT CATGTCCCOT TCC CTATAGACCG TTAATGTTAA AAGGGTCTCO TTAACTAAAA GTACAGGGCA AGG

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#### FIGURE 45A

	10	20	30	40	50	6:
	GATGCTATTT CTACGATAAA	GGGCAATTTC CCCGTTAAAG	TTATTGACAG AATAACTGTC	TTTTGAAATG	TTAGGCTTTT AATCCGAAAA	ATCTCCATTT TAGAGGTAAA
61	TTTAGTACTT	AAATTTTCCA	ACATGGGTGT	TGCTTGTTAT	TTTATCAGTA	TAAAATAGAA
	AAATCATGAA	TTTAAAAGGT	TGTACCCACA	ACGAACAATA	AAATAGTCAT	ATTTTATCTT
121	GAGTGGTTCT	GTTCTGGAAT	TTAGTATATA	CATGAGTATC	TAGTGTATGT	CAGCCATGAA
	CTCACCAAGA	CAAGACCTTA	AATCATATAT	GTACTCATAG	ATCACATACA	GTCGGTACTT
181	AATGAACCTT	TCAGATGTTT	AACTTCAGGG	AACCTAATTG	AGTCATTGCT	CCAGACATTG
	TTACTTGGAA	AGTCTACAAA	TTGAAGTCCC	TTGGATTAAC	TCAGTAACGA	GGTCTGTAAC
241	TTGCTTTGAA AACGAAACTT	CCCACTATAT GGGTGATATA	TENNERNANCE AMERICAN SERVICE AMERICAN SE	CGGGCAATER GCCCGTTACT	CTCAGTGTGG GAGTCACACC	CAAGGATACT GTTCCTATGA
301	ACTGCAGGCC	TGTTTCTGGA	AGGCACTGGA	STOCTOTGAT	GCAAACTTTG	GCCAGGGACT
	TGACGTCCGG	ACAAAGAECT	TCCSTGACCT	SAGGAGACTA	CGTTTGAAAC	CGGTCCCTGA
361	CCTTGATAGC	TCTTAAATAG	ATGCTGCACC	AACACTOTOT	TTCTTTTCTC	TCTTTTCTT
	GGAACTATCG	AGAATTTATO	TACGACGTGG	TTGTGAGAGA	AAGAAAAGAG	AGAAAAASAA
421	TATTCAATAT	TAGACTACAA	GCATTITAAI	GACTTOTOAG	GGTTTCTAGC	TCTCTCTCAT
	ATAAGTTATA	ATCTGATGTT	CGTIAGATTI	STGAAGAGTO	CCAAAJATCG	AGAGAGAGTA
48:	TTCACACATG	CTTTICTAGT	AATCTCTACT	CATATATCTT	ACTGCTACGC	TGGGGCCAGA
	AAGTGTGTAC	GAAAGGATCA	TTAGASATGA	GTATATAGAA	TGAGGATGCG	ACCCCGGTCT
541	TAACHHHNHH	CTTCCATTTT	GTTTTTATCT	CTATTCTTCT	TCCCCTTCTG	CTTTCATTAT
	ATTGHNHNHH	GAAGGTAAAA	CAAAAATAGA	GATAAGAAGA	AGGGGAAGAC	GAAAGTAATA
601	TGAAACTTTC ACTTTGAAAG	TGCTTTCATT ACGAAAGTAA	ATTGAAACTT TAACTTTGAA	TCCCAGATTT AGGGTCTAAA		ACCTGGCATT TGGACCGTAA
661	GGAACTGTTT CCTTGACAAA	CCTCTTCCCT GGAGAAGGGA	GTGCTGCTTT CACGACGAAA	CTCCCATTGC GAGGGTAACG	CATGTCCTTT GTACAGGAAA	TTTTTTTTT
721	TTTTTTTTTT	TGAGACAGTG	TCACTCTGTT	GCCCAGGCTG	GAGTGCAATG	GTGCAATCTT
	AAAAAAAAA	ACTCTGTCAC	AGTGAGACAA	CGGGTCCGAC	CTCACGTTAC	CACGTTAGAA

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#### FIGURE 45B

781	GGCCACTGCA	ACCCCGACTC	CGGGTTCAAG	TGATTCTCTA	CCTGCCTCAG	CCTCCTGAGT
	CCGGTGACGT	TGGGGCTGAG	GCCCAAGTTC	ACTAAGAGAT	GGACGGAGTC	GGAGGACTCA
		•				
841	AGCTGGGATT	ACAGGTGCCA	CCACTATSCC	GGCTGATTTT	STATTTTAGT	AGAGATGGGT
	TOGACCOTAA	TGTCCACGGT	GGTGATACGG	CCGACTAAAA	CATALAATCA	TETETACCEA
901	TCACATGCAG	ATCAGCTGTT	CCGACTCTGA	CCAGNIHINNN	инининини	ATCAAAGTCA
		TAGTCGACAA				
٠ ; :	GCCAAAGTGC					
	COSTTTCACS	ATCCGAATCT	CATTANCACA	TTAAAGGTGT	GTTCACGTTG	GATCACATTA
1:::	SCCTCAAGAA	TGTNNNTATG	AATGTCTCGA	ACGTTAGTAA	CTANTANCAN	GTAGTTAGTT
	COGAGTICTI	ACADODATAC	TTACAGAGCT	TUCAATCATT	GATTATTGTT	CATCAATCA
1681	TATAGATGTA	TOOTAGTATO	TAGCA			
	1717071017					

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### FIGURE 46A

	10	20	30	40	50	60
1	CACAAAAAA GTGTTTTTT	GATTATTAGC CTAATAATCG	CACAAAAAA GTGTTTTTT	CCTTGAAGTA GGAACTTCAT	ACGCATTAAA TGCGTAATTT	ATGTTAATGG TACAATTACC
61	ATTCACTTTA TAAGTGAAAT	TTGAGCATCT AACTCGTAGA				
121	ATACGTCATT TATGCAGTAA	TAAACCTTAC ATTTGGAATG				
181	GCACAGGAGG CGTGTCCTCC	CTTAGATAAC GAATCTATTG				
241	ATTCAAATTA TAAGTTTAAT	TTGATAAGAA AACTATTCTT				
301	CGCTTTCCAG GCGAAAGGTC	AGCATGTGCT TCGTACACGA	GTTGATAGAG CAACTATCTC	CTTGATGTCT GAACTACAGA	AACTCTCTGA TTGAGAGACT	AATTTTCCAT TTAAAAGGTA
361	TCTTATTTGT AGAATAAACA	CTCACTGGTA GAGTGACCAT				
421	ACAGGAGGAT TGTCCTCCTA	CAAAGATAGG STTTCTATCC	ATTTCATTTA TAAAGTAAAT	SAATSOCTAA CTTAGGGATT	AGCTTCACGT TOGAAGTGCA	ATTTTAATTC TAAAATTAAG
481	AGAATAAGAT TOTTATTOTA	TCAGGCAGAC AGTCCGTCTG	CACCAGTATA GTGGTCATAT	TECCATEGIC ACCGTACCAG	CCTGGTTATC GGACCAATAG	TTTCAGCAGG AAAGTCGTCC
541	TGACCGAGAA ACTGGCTCTT	AGAAAACATG TCTTTTGTAC				
601						ATGGGCATGT TACCCGTACA
661						GACAAACTTA CTGTTTGAAT
721	AGAGACTGAG TCTCTGACTC	TGTTCAAACT ACAAGTTTGA	GAATAATCTC CTTATTAGAG	GACCTTAATT CTGGAATTAA	GTAACTATAT CATTGATATA	TTTATGAAAT

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#### FIGURE 46B

- 781 CCAGCTGTAA GGCAAAACAG ACTOTTGGCT ACACGGCATT TGTCTGTTAA TGATACTCAA GGTCGACATT CCGTTTTGTC TGAGAACCGA TGTGCCGTAA ACAGACAATT ACTATGAGTT
- 841 COTTAACCGT CACTTAATAA TGCTGAATAA TGTCATTAAT CTGAGATGTT AGTATGATCA GGAATTGGCA GTGAATTATT ACGACTTATT ACAGTAATTA GACTCTACAA TCATACTAGT
- ### ATGGEARTER CTGCTGAGCT CTGGRAGCCC TACCCTTAGT GACGACTCGA GAGCTTCGGG

# FIGURE 47A

			0.0			
-239 -120 -1	2 2	160	270	360	450	340
E C S	53	84	Ë Ĉ	85	25 C	CCT TIC AGI GCT TIC TCT CCT CAA GCA ATG CCA GAG GCC GAT CTA GTG TAT GTT AAC TAT GCA Pro Phe Ser Ale Phe Ser Pro Gin Gly Het. Pro Glu Gly Aep Leu Vel Tyr Vel Aeb Tyr Ale
XXXXXXXXX	84	₹;	<b>A 6</b>	TAC Tyr	g £	32
2000	63	A T	35	Ser.	25.5	¥ \$
261 CTCAAAAOOOOCOCATTTOCTT CTGGAAOGAATTOCAOCCTGCAAOOCT SOOGAAOCOOCTCTGCTGOOOCOAA	TOO OUT GTG ONC ACC UND CON CON CON TOO CTG TOO OCT DOG OND CTG GTG CTG OND OCT Set Ale Val Ale The Ale Aig Arg Pro Arg Irp Leu Cye Ale Gly Ale Leu Val Leu Ale Gly Ale Leu Val Leu Ale Gly Ale Leu Val Leu Ale Gly	GUS TOS TIT ATA AAA TOO TOO AAT GAA GUT ACT AAC ATT ACT COA AAG CAT AAT ATG AAA GLY TEP IN 110 110 110 110 100 Set Aen Glu Ale Int Aen Ile The Pro Lye Bie Aen Het Lye	ATC AND AND TIC ITA TAT ANT TIT ACA CA: ATA CCA CAT ITA GCA GGA ACA GAA CAA AND ITO Lye Lye Lye lie Leu Tyr Agn The Glin Ite Pro His Leu Ala Gly The Glu Glb Asn	TOG AAA GAA III GOC CIG GAI ICI GII GAG CIA GCA CAI IAI GAI GIC CIG IIG IAC	ATA ATE AAT GAA GAI GGA AAT GAG ATT TIC AAC ACA TCA ITA ITI GAA CCA CCI CCT CCA II. II. Ann Glu Ann Glu II.	GTT Vel
SAAT	23	CAT B1.	ĄĘ	CTG Leu	អ្ន	Σž
C PC	84	<b>№</b>	017	CTC Vel	35	610 V
261 7010 2000	85	85	AL.	GAT A.P	E E	ניא
250	S &	ACT	IIA	TAT Tyr	114	A ST
200	5 S	ATT 11.	CAT FILE	7 <b>.</b>	Ž.	85.7
CT0C/	23	¥ \$	y :	¥ 4.	A E	350
YVV KX.Y	53	ACT 11.r	ATA 11.	CTA Leu	¥ V V	ប្តូរ
7 P.	₹ 8	RI.	֓֞֟֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֟֓֓֓֟֓֓֓֟֓֓֓֟֓֓֓֟֓֓֓֟֓֓֓	GAG	7 5 •	ATG.
).TOG	OCC Pro	₹; ;	ACA	CT GTT	T .	8 <del>2</del>
EX.1	Ar &	A	ËĒ	Ser	2000	<b>₹</b> 5
CAC)	00; A18	7€. Ser	AAT Aga	CAT A.p	<b>A</b>	CC1
<b>₹</b>	AL.	155 166	TAT Tye	CTG L.	66A 61y	17.1 Ser
1101	A t	A.A.A. 1.y.	TTA Leu	85	₹ \$ £	710
A. 16	Ate	¥=	TTC In.	111 51.0	<b>₹</b> 200	₽¥
7.A.C.	24.2	Es	AAG Ly•	<b>₹</b>	AAT Aen	AGT Ser
Taxa Seec	Ale Ve	23	AAG 1. y •	₹\$	ATT -11•	77.
XXAT.						
XTX	\$ &	E E	¥ ¥	CAG	17. 3.	S S
7. T. T. C. A. C.	P F	55.3	CAG G14	30.5	CAT CCC AAC TAC ATC Bie Pro Aen Tyr Ile	C1A V•1
75.75	3 2	5 E	A1.	<b>₹</b> 5	175	A17
AGA1	3:	327	₹\$	ATT 11.	\$ <b>\$</b>	CAT
51.75 51.75	11.	53	23	<b>₹</b> 5	85	100 Ser
AGAT	23	53	33	<b>1,</b>	CAT B1	C11
A00C 6400	<b>44</b>	Eė	SAT A B B	84	Act	AAT
261 CTCAAAAAAAAAAAATTTOCTT CTCCTOGAACCAAATTTOCCTCTCTCTCTCTCTCTCTCTCTCTCTCT	ATO TOO AAT CTC CTT CAC GAA ACC Het Trp Aan Leu Leu Bie Glu Thr	GLY Phe Phe Leu Leu Gly Phe Leu Phe	THE ING GAT GAA TTG AAA CCT GAG AAC Phe Leu Asp Glu Leu Lys Ale Glu Asn	CAG CTT CCA AAG CAA ATT CAA TCC GLB Leu Ala Lya GLb Ile GLb Ser	AAT AAG ACT C	TAT GAA AAT GIT ICG GAT AIT GIA
63	45	89	Eě	36	TA PA	TAT TY
				•		

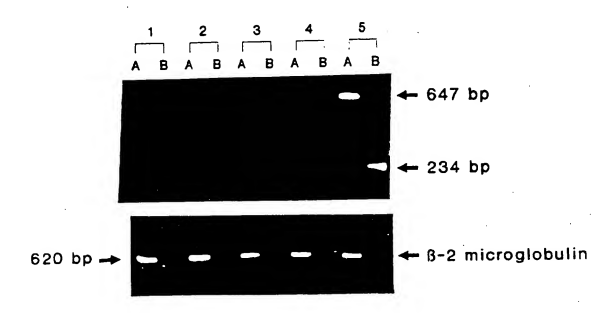
# FIGURE 47B

630	720	810 270	300	990	1080
Y S	AAG Lye	ភ្ជ <b>ះ</b>	TAT	25 to	58
5.5	010	Q E	TAC	CTT V•1	ATA
G11 V•1	35 17 17	CTC Lea	ž÷		200
Ly.	Pra		11.	37.	M.
930	SCI Ale	CAC	Pro Pro		IAC
1AT 17E	E É	GCA GLy	CAT	GTG V-1	ATT
AGA Arg	1AC 1y1	 	CIT Vel	₹;	V P
¥ 8 €	GAC A * p	61.7	CCT Pro	C1C Leu	ACA The
GTA ATT Val 110	A1.	¥ ¥	A11	AUT Ser	GTG V
GTA Val	Rts	CTG Leu	AGT Ser	617	۲ کا تا کا
A17 11•	CAC Asp	AAT CTG	ž ŝ	۸:۸ ۲:۸	AAT
Ly.	3.5	C17 L•	CTT E	16; 1rp	N €
35	17C	ATC 11•	COT CIT	X	֭֭֭֭֭֭֭֭֭֭֭֓֞֞֝֞֜֞֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓
1CT Ser	= -	¥ ¥	C.T.T VA.I	AINT AINT TOTAL	Ž =
AAT TISC A	CHE ATT CTO	6.1.y	AL VAL	CAT A P	ATS CAC ATC CAS ICT. ASS AAT GAA GIG ACA AGA ATT IAC AAT GIG ATA OCT. Het His 116 His See The Ass Glo Val The Ass Fig. The Ass
AAT.	7 × ×	CGT Ara	3 5	1.1.A	E AG
AAA A16 1.ye 11e	4 T	: AG	Ω.V VI•	CCA Fra	A15
	AAA - 7	61C V•1	25A ATT (1)	A 14	GTC AAG
A1G Het	\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$	5 5	A	30.CA	CTC
1 Ans	500	1 GET	ACK; CGT	0.17	1 AA 1
Ara	AX.	C. C. A.			35
₹ 0 C C C C C C C C C C C C C C C C C C		r cct	TAT Tyr	A ATG	ACA T
110	OCC CAG (TG	1101	r ocr	A AAA 1.7*	7 TCT
1 kk	1 000 E	G AAT	A TAT	A GAA	THE ST
7. TT . 2	GTT ANA AAT	T TGG 7 Trp	T GAA	C CTA	A AMC
GAC TTC Asp Pho Intron	₩ **	T 001 P 617	A AAT	6 CTC	1 0CA
\$ <b>2</b>	10 g	A GAT	A OCA	G AAG n Ly•	TACT
r GA	AAT AAG Aen Lye	r Pro	C CCA	A CAG	TIT N
A ACT		X 1AI	IT TAC	P Ale	1 000 140 140
48	45	B.	95 14 14	GAT	85

# FIGURE 47C

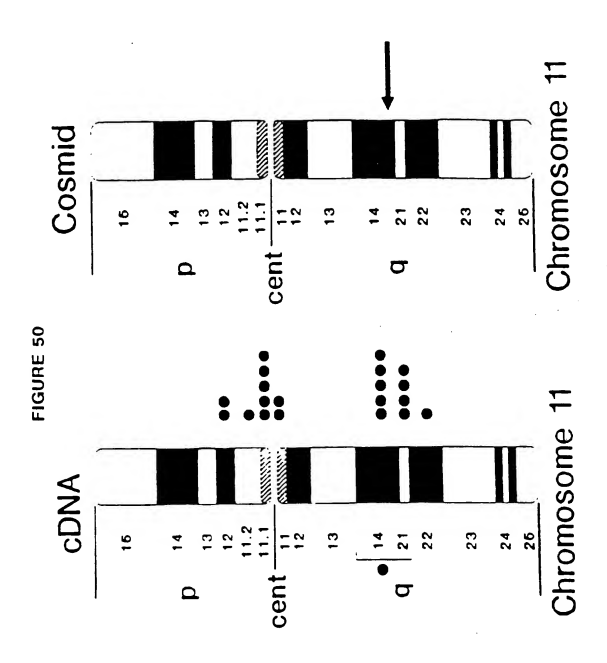
75/130								
38	1260	1350	1440	1530	1620	1710 370		
Gly Ile Asp Pro Gln Ser 390	AOC Ser	ATTA 111•	046 614	85	MA S	Eė		
Gln	\$ <del>*</del>	TAT Tyc	<b>₹</b>	ATG Het	1:8	ATG		
Pro	E &	AL.	ŽĘ	617	A T	8 <del>2</del> 8		
\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	23	GT0 Ve.1	1. 1. 1.	AGT Ser	IXI,	48		
11.	114	95 91,4	¥¥ V*t¥	5 E	¥8	TAT Tyr		
213	P P	Are FGT	3 =	GAG	84	E É		
617	AGA Are	CVO	GTA V•1	Pro	ACA	AAG Lye		
==	ACA Arg	¥5	23	TCC Ser	85.	85		
0 T	CCT Pro	CTT	AGC Ser	oct Pro	\$ 5.5 \$ 5.5	CTG Vel		
rec cre	AGA Are	AGA CTC CTT CAA	TAC	AGT Ser	A 4.	TTG Leu		
<b>5</b>	8 r	AGA Ar &	ATG Het	AAA Lye	A11	GAG		
Arg Any Ser	623 634	S CA	CTG Leu	Ly.	\$ t 5	TAT		
y s	GAA GLu	AAT A B B	g :	D.E.	E3	A F		
E	AAG L.y.	Glu Aen	A:A Tir	15 7 - 7	Are Are	CAA Clu		
GGT CAC	₹.	35	C, 51	ACT Sec	35	TAT Tyr		
4 7	010 [.•]	GAG TOS G'A Glu Trp Ale	\$ £	<b>₹</b>	110 710	GTC Val		
1 2	ACA The	1. E	C111 V•1	1A1 17.	EE	ACT Ser		
AIT CTG	C 1.7	SA:	AGA Ara	CTT Leu	CTC V•1	3=		
TAT GTC /	E É	T. F.	ACT CTG The Lou	TCT Ser	35	CTG TAT		
TAT 175	AUC Ser	17.	ACT The	¥:	E.E	CTC Lev		
ξ¥ V	A: 8	5.17	1AC 171	25	CAT A.P	ង្គ		
ئے کو کو کو کو	CTC ACG	GI CII CII GII Gly Leu Leu Sly	AAC	\$ ? 0 ?	AAT Aen	TAT		
§ 🗜	GAA ATT	110	\$ <del>1</del> 5	60C 111 617 Ph•	8 t	85.4		
3 :	5. 1.3 1.3 1.3 1.3 1.3 1.3 1.3 1.3 1.3 1.	25.5	₹°	600 617	7CT 5.ec	S. S.		
CT0 -	3=	Eé	ATA 11.	<b>61.4</b>	5 5	₽.€		
84	715 V•1	33	101 Ser	A.P	13			
\$ 6	GTT GTT	CAA GAA	TCA Ser	Pro Pro		\$ <b>Q</b>		
£ §	84	84	CAC	AOC Ser	Age a	ĄĘ		
£3	84	A P	A.	¥.	ATA 1			
ğ	449	<b>B</b> =	<b>A</b>	63	r g	8 5		

#### FIGURE 48



# 

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#### FIGURE 51

8 9 M H 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 X Y

••

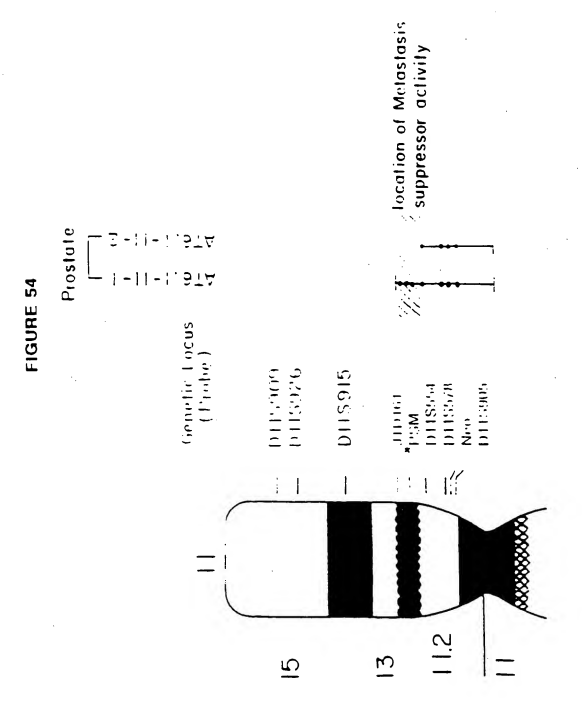
# FIGURE 52

	•					clone 1	clone 2				1 clone 4	1 clone
Markers	Uncut	RNA	LnCap	PC3	AT6.1	AT6.1-11	AT6.1-11	۷۵	A9 (11)	R1564	R1564-11	R1564-11

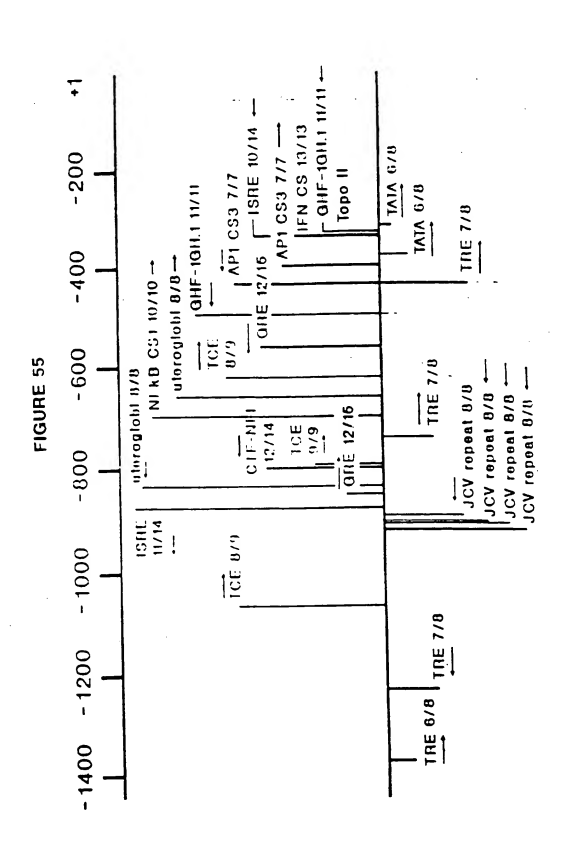
# FIGURE 53

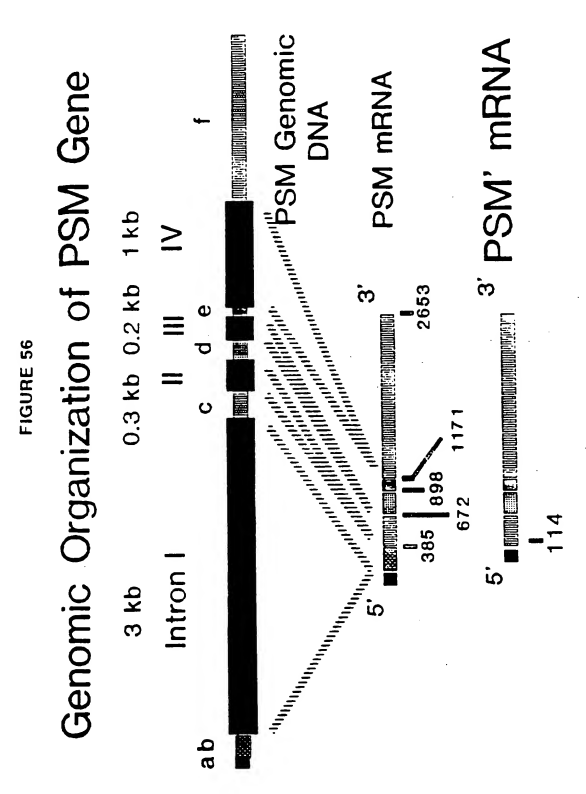
# VNN INST INSU DNI ADI NOCARCINOMA LOCULAR DOCACA RALPROSTATIC RAT MAMMARY THURLDY VINODAINSOMAL MOUSE 1111 7 HUMAN MAMMARY HUMAN PROSLAM TISSUE/ CELT R1564-11-C1.5 R1564-11-C1.6 AT6.1-11-CL2 R1564-11-C1.2 AT6.1-11-CL1 R1564-11-CT.1 10(11) K1504

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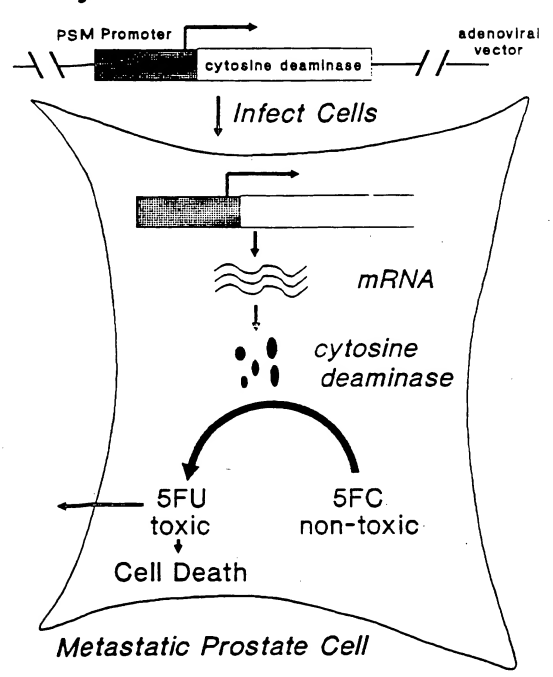


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Prostate Specific Promoter:
Cytosine Deaminase Chimera



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#### FIGURE 58A

	10	20	30	40	50	εc
:	GCGCCTTAAA CGCGGAATTT	AAAAAAAAA C TITTTTTTT A	TTCTTGGAÅ AGAACCTT	AATGTCCAGC TTACAGGTCG	TCTTGCTTAA AGAACGAATT	TAAAAATATATATATATATATATATATATATATATATA
é l	GAAAGGAAGA	AAGAGACTCT C	CTCTCTCCA	CTCCTATAAT	TATGAGGAAC	TTTTATTCAA
	CTTTCCTTCT	TTCTCTGAGA G	GAGAGAGGT	GAGGATATTA	ATACTCCTTG	AAAATAAGTT
121	CTITGAAATT	CTATACAATC T	CTACAATAC	TOTACTGAAT	AAAAGCAGAG	CASAAAAGC
	GAGACTITAA	GATATGTTAG A	GATGTTATG	AGATGACTTA	TTTTCGTCTC	GTCTTTTTCG
181	TGCGCTTTTT ACGCGAAAAA	TTCCATAGTC C	TTOETAADDE AADDATTOOL	GTCATCAGTG CAGTAGTCAC	TAAATCACCA ATTTAGTGGT	CCGCGCCTT GGCGCGGGAA
241	TTTCCTAAAG AAA33ATTT0	AATATTATTG :	AATAATAA TTATTAATAA	ACATGTAGGG TGTACATCCC	TATTATCCTC ATAATAGGAG	CACTTACATT GTGAATGTAA
301	ACAAAACCAT	TTTTTAAAGC	CGGGGGTGGT	CODORATORS	TSTAATSCCA	GCACTTTGGG
	TGTTTTGGTA	AAAAATTTCG	GCCGGGAGGA	DESERVED	ACATTAGGGT	CGTGAAACCC
361	AGGCCCAGAC	AGGIGGATCA	CCAACTCGAG	AAATCGAGAC	CATCCTGGCC	AACATGGTGA
	TCCGGGTCTG	TOCGUITAGT	CCTTCAGCTC	TITAGCTCTG	GTAGGACCGG	TTGTACCACT
423	AACCCCATCT	CTACTAAAA	TA DAMAAATI	PODDODÁBIL	GSTGGCGGGG	TOCTGTAGTS
	TTGGGGTAGA	GATGATTTT	AT STOTTTAA	KODODÁKIT	A COACCGCOC	AGGACATSAG
4 8 3	1 CONSCINCT	ASSASSOTSA	000A00A0A0A0	A TOSCTTGAAC	COGGGAGGCC	GAGGTTGCAG
	GGTCGATTA	S TOSTOSTACT	000T:0T:T	TASCGAACTT	G GCCCCTCCG	CCTCCAACGTC
54	1 TCAGCCAAG	A TAGOGOCAGT	SCACTGGAS	T CTEGTGACA	G AGTGAGACT	C CCTTAAGAAA
	AGTCGGTTC	T ATCGOGGTGA	CSTGACCTC	E GACCACTGT	C TCACTCTGA	G GGAJTTOTTT
60	1 GAAASSAAG	G GAAGGGAAAG	SGAASSAAS	S SGAGGGGAA	S GGAGGGGAG	G GGAGGGGAGG
	CTTTCSTTC	C CTTCCCTTTC	COTTSSTTC	C CCTCCCCTT	C CCTCCCCTC	C CCTCCCCTCC
66	1 AAAGAAAAG TTTCTTTTC	A ATACTGGAAC T TATGACCTTG	TTGTTGAAG AACAACTTO	G CAGAGACTT C GTCTCTGAA	T ATTTTCATA A TAAAAGTAT	T CCCGGCTATG A GGGCCGATAC
72	1 TCTGGCTAC AGACCGATG	T GTCTTACGTA A CAGAATGCAT	ATAGATATA TATCTATAT	A AATCAATCI TTAGTTAGA	T GGTTGGAT A CCAACCTAI	TA ACCAGAAGAA AT TGGTCTTCTT

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#### FIGURE 58B

781	TGAGAAGATA ACTCTTCTAT	TATTCTGGTA ATAAGACCAT	AGTTGAATAC TCAACTTATG	TIAGCACCCA ( AAICGTOGGT	CCCCATTAGT	GCTTGGACAG CGAACCTGTC
841	GACCAGGTCC CTGGTCCAGG	AAAGACTGTT TTTCTGACAA	AAGAGTCTTC TTCTCAGAAG	TGACTCCAAA ACTGAGGTTT	CTCAGTOCTC GAGTCACGAG	CCTCCAGTGC GGAGGTCACG
901	CACAAGCAAA GTGTTCGTTT	CTCCATALAG GAGGIATITC	GTATCETGTG CATAGGACAC	CTGAATAGAG GACTTATCTC	actgtagagt Tgacatctca	GGTACAAAGT CCATGTTTCA
961	AAGACAGACA TICTGTCTGT	TTATATTAAG AATATAATTC	TCTTAGCTTT AGAATCGAAA	GTGACTTCGA CACTGAAGCT	ATGACTTACC TACTGAATGG	TAATCTAGCT ATTAGATCGA
1021	AAATTTCAGT TTTAAAGTCA	TTTACCATGT AAATGGTACA	GTAAATCAGG CAITTAGTCC	AAGAGTAATA TTCTCATTAT	CAACAAACCT CTTCTTTOGA	TGAAGGGTCC ACTTCCCAGG
1081	CAATGGTGAT GTTACCACTA	TANATGAGGT ATTTACTCCA	GATGTACATA CTACATGTAT	ACATGCATCA TGTACGTAGT	CTCATAATAA GAGTATTATT	GTGCTCTTTA CACGAGAAAT
1141	AATATTAGTC TTATAATCAG	ACTATTATTA IGATAATAAT	GCCATCTCTG CGGTAGAGAC	ATTAGATTTG TAATCTAAAC	ACARTAGGAA TGTTATCCTT	CATTAGGAAA GTAATCCTTT
1201	GATATAGTAC CTATATCATG	ATTCAGGATT TAAGTCCTAA	TTGTTAGANA AACAATCITI	GAGATGAASA CTCTACTICT	AATTCCCTTC TTAAGGGAAG	CTTCCTGCCC
1261	TAGGTCATCT ATCCAGTAGA	AGGASTTGTC TCCTCAACAS	ATGGTTCATT TACCAAGTAA	GTTGACAAAT CAACTGTTTA	TAATTTTCCC ATTAAAAGGO	AAATITTTCA TITAAAAAA
1321	CTTTGCTCAG GAAACGAGTC	AAAGTCTACA TTTCAGATGT	TOGANGENCO AGETTEGTGS	CAAGACTGTA GITCTGACAT	CAATCTAGTO GTTAGATCAG	CATCTTTTC GTAGAAAAAG
7367	CACTTAACTC GTGAATTGAG	ATACTSTGCT TATGACACGA	CTCCCTTTCT GAGGGAAAGA	CAAAGCAAAC GTTTCGTTTG	TGTTTGCTAT	TCCTTGAATA AGGAACTTAT
1441	CACTOTGAGT GTGAGACTCA	TTTCTGCCTT	16101ACTCA ACGGATGAGT	SCTGGCCAT CBACCGGSTA	COGG GOATTI	CALAGAAGAG
1501	ATCTCCACTG TAGAGGTGAC	GGTCANATCC	TACCTGTACG ATGGACATGG	TTATESTICT	CANTTITOG	CACGAAGGTA
1561	AMSTACTOC	TAGCAAATGC	ACGSCCTCTC	TOACGGATIA	TAASAACAC	A GTTTATTITA
	TTTCATCACG	ATCGTTTACG	TSCCSGAGAG	AGTGCCTAAT	ATTCTTGTG	CANTANAT
1621	TARRECATGT ATTTCGTACA	AGCTATTCTC TCGATAAGAG	TCCCTCGAAA	TACGATTATT ATGCTAATAA	ATTATTANG	A ATTTATAGCA I TAAATATCGT
1681	GGGATATAA? CCCTATATIA	TTTOTATGAT AAACATACTA	GATTOTTOTO CTANGANGA	GTTANTOCAL CANTIAGGTT	CCAAGATTG GGTTCTAAC	A TTTTATATCT T AAAATATAGA
	TARTGCATTO	- Torextees	Cidinicas			C TOCCTTCAAC
	TICHOGIC	A TANGAMAN	A AAGOAGGG			T CCCCTTCCTT
186	COCTITICCE	T TCCCTTCCT	TOTTTCTTG	A GGGAGTETC T CCCTCAGAS	A CTCTGTCA T GAGACAGT	CC AGGCTCCAGT OG TCCGAGGTCA

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#### FIGURE 58C

1921	GCAGTGGCGC CGTCACCGCG	TATCTTGGCT ATAGAACCGA	GACTGCAACC CTGACGTTGG	YOCCEGYOCC	CGGTTCAAGC GCCAAGTTCG	CTANGAGGAC
1981	CCTCAGCCTC GGAGTCGGAG	CTGAGTAGCT GACTCATCGA	GGGACTACAG CCCTGATGTC	CAGCCCCCA CTCGGGCGGT	CCACGCCCAG GGTGCGOGTC	CTARTITITIC GRTTANAC
2041	TATTTTTAGT ATAAAAATCA	AGAGATGGGG TCTCTACCCC	TTTCACCATG AAAGIGGTAC	TTGGCCAGGA AACCGGTCCT	TGGTCTCGAT ACCAGAGCTA	TTCTCGACTT AAGAGCTGAA
2101	CGTGATCCGC GCACTAGGCG	CTGTCTOGGC GACAGACCCG	CTCCCAAAGT GAOGGTTTCA	GCTOGGATTA CGACCCTAAT	CAGGCGTGAG GTCCGCACTC	CCACCACGCC GGTGGTGCGG
2161	CCCCANATITY CCCCANATITY	AANGGTTIT TTTACCAAAA	GTAATGTAAG CATTACATTC	TGGAGGATAA ACCTCCTATT	TACCCTACAT ATGGGATGTA	GTTTATTAAT CAAATAATTA
2221	AACAATAATA TIGTTATTAT	TI CTTTAGGA AAGAAATCCT	AAAAOOOCGC TTTTCCCGCC	GGTGGTGATT CCACCACTAA	TACACTGATG ATGTGACTAC	ACAAGCATTC TOTTCGTAAG
2281	CCGACTATGG GGCTGATACC	AAAAAAGCG TTTTTTTCGC	CASCTTTTTC GTCGAAAAG	TGCTCTGCTT ACGAGACGAA	TTATTCAGTA AATAAGTCAT	GAGTATIGIA CTCATAACAT
2341	GAGATTGTAT CTCTAACATA	AGAATTTCAG TCTTAAAGTC	AGTIGAATAA TCAACTTATT	AAGTTCETCA TTCAAGGAGT	TAATTATAGG ATTAATATCC	AGTGGAGAGA TCACCTCTCT
2401					GAGCTGGACA CTCGACCTOT	
2461					TTTCCTTCTC ALAGGALGAG	
2521					CTAGAAACAC	
2581					AGGGCTGATA TCCCGACTAT	
2641					CYCCCCCCCC	
2701					CCGAGATGTG GGCTCTACAC	
2761					GCTGGCTGTG CGACCGACAC	
2821					TCOOTAGGG AGCCATCCCC	
2881					CCCTGAGCGC	GGTCAGCTGC CCAGTCGACG
2941						GGTAATOTGG CCATTACACC
3001	OOTGAGCACC CCACTCOTGG					

Acividin

Azotomycin, becomes active by in vivo conversion to DON

6-diazo-5-oxo-norleucine, DON

HOA .

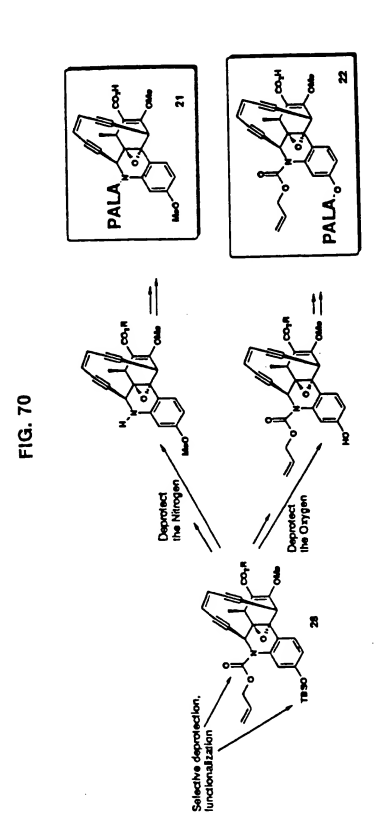
FIG. 65

FIG. 65

FIG. 65

$$N_2$$
 $N_2$ 
 $N_3$ 
 $N_4$ 
 $N_2$ 
 $N_4$ 
 $N$ 

FIG. 6



# FIG. 72A

09.	GTGCTGGGAC CACGACCCTG	GGCGATTAGO	GGAACGGTGC CCTTGCCACG	gtagaactga catcttgact	GACAGAGGAA CTGTCTCCTT	TGTTTGTTTG	GCTTGGGAAC CGAACCCTTG	<b>GCTGITITITC</b> CGACAAAAG	ACACAGGCAA TGTGTCCGTT	GCCTTGAACA CGGAACTTGT
80	TGGTGCCGCG (	CCCCAGGGGC	AGGGTAGCTG TCCCATCGAC	CAGGTTGAGG GTCCAACTCC	AGCCCTGCAA TCGGGACGTT	TTGTTTTGTT AACAAAACAA	ACAGAGGCAA TGTCTCCGTT	CGGGTCTTTT GCCCAGAAAA	AAGCAGAACC TTCGTCTTGG	CTTCTTAGTG Gaagaatcac
0-	GTCTTCCCCG CAGAAGGGGC	TGCTGGTCTT ACGACCAGAA	acttaggagg Tcaatcctcc	GACAGTCACT CTGTCAGTGA	CAAGTGCTGG GTTCACGACC	TTGTTTTGTT AACAAAACAA	CTTGGAAGTA GAACCTTCAT	TCTTTACCAG AGAAATGGTC	TITCTAAGAA MAGAITCTT	gactitgcca Ctgaaacggt
CE .	AAACCTCGGA TTTGGAGCCT	GGGATCCTGT CCCTAGGACA	GCACCCCTCG	CTGCTGGTAG	AGGAAGGTTC TCCTTCCAAG	TTGTTTTGTT AACAAAACAA	TTCTTTCTTC AAGAAAGAAG	TCTGGACAGG Agacctgtcc	ttgatccaac aactaggttg	ttccagtttt Aaggtcaaaa
20	CCTCGCGGAG	GCTGCCGAGT	TGTGGGGTGA Acaccccact	TCTC3ACAAG Agagctgttc	AACTGGGCGT TTGACCCGCA	TGCTTTTGTT ACGAAAACAA	TCTCTGTGCA AGAGACACGT	TGTGTGAACC AGGTCAGCAA ACACACTTGG TCCAGTCGTT	attigcagac Taaacgicig	TTTTATTAAA AAAATAATTT
10	TAGGGGGGGG	TCGCGGGTCA	GTCGGGGTAA CAGCCCCATT	AGGGCTGAGT TCCCGACTCA	GAGAACCTGA	GITTITITI CAAAAAAAA	TTTTTTACC AAAAAATGG	TGTGTGAACC ACACACTTGG	CTGGGTACTG Gacccatgac	GCTCAGACTC
	ન	19	121	181	241	301	361	121	481	541

ATATTATCTG GGATAAAATA CTACTCCTAT TATAATAGAC GATGAGGATA CCTATTTAT CGTTAGTTAC AGGGAGAGTC GCAATCAATG TCCCTCTCAG AGTTACCGAG TCAATGGCTC 601

TACTGGGATT ATGACCCTAA CCTAGCACAG TACATTTAGA ATGTAAATCT ATAATATAGC TATTATATCC GTAATACTAA CATTATCATT GTTTAATAAC CAAATTATTO

661

TACACAGGAC ATGTGTCCTG TACTCCTCAT TGGACTTTAA ATGAGGAGTA ACCTGAAATT ATANAGAAGA AAATGGTTCT TTTACCAAGA TATTTCTTCT AAGCGGTGAA TTCGCCACTT 721

TCTTTCGGGA GACGAGOCTT AAGAACTGGG AGAAAGCCCT TTCTTGACCC CTCCTCGGAA TATCACCAGG TAGTCCACTC ATAGTGGTCC ATCAGGTGAG TAGTCTAAGG ATCAGATTCC

781

TAGGGCATGG ACCAGATGGG TITAAACAAA TICAATATCT TCCACTAGGT ATCCCGTACC TGGTCTACCC AAATTTGTTT AAGTTATAGA AGGTGATCGA GATITITIDAA CCACACACTG IGCICATAAC AAICTICAIC TTTAGAAGAA AAATCTTCTT 841

GTTGTTAAAA

TCACCTTGGG

106

TTAGAAGTAG ATTCCGTGCT GGTGTGTGAC ACGACTATTG ACCETCACTC TCATCCCTGT CTAMANACTT CAACAATTTT AGTGGAACCC

TAAGGCACGA TAAAATAAGA AGGACCATAA CGGGAGTGAG AGTAGGGACA TCCTGGTATT ATTTTATICT TCTTAAAAGG AGANTTTTCC 961

1081 GCTCTCCCCT CCCCCTACAG GCCTCCATCC TCTTCATCCT GTTCATITT CAGAICTCAG	CERERAGER GEGGERTOTO COCRECINGO AGARCINGGA CARGINAAAA GICINGAGIC
GIICAILITI	CAACTAAAA
TCTTCATCCT	AGAAGTAGGA
GCCTCCATCC	CGGAGGTAGG
CCCCCTACAG	したさんないかいかい
GCTCTCCCCT	なりかりかんかなかい
1081	

	1141 TTCAAGCATC TCGTCCTCAG TGTGGTGTTT CCTGATCCCT CACTCTAATC CAAGTCTTTC AAGTTCGTAG AGCAGGAGTC ACACCACAAA GGACTAGGGA GTGAGATTAG GTTCAGAAAG
CGAGAGGGGA GGGGANIGIC CGGAGGINGC NGMGGMGGM GMGGGGAGGA	CACTCTAATC GTGAGATTAG
	CCTGATCCCT GGACTAGGGA
200000000000000000000000000000000000000	TGTGGTGTTT ACACCACAAA
212122222	TCGTCCTCAG AGCAGGAGTC
ところうらんのとうと	TTCAAGCATC AAGTTCGTAG
	1141

CUBARA MARIE BARRARA COMPOSITION OF THE COMPOSITION	201414111	じしなけななけるべる	ACAAAATACG TGTCCACCTT AGAATAAAGG CAAACGCAGG 11AG1ACA1A ACAITA1ACG
FAFFACEA	141010115	TEXT KEUKEE	<b>ビオケンドイラビイエ</b>
	ンフィランライエーラ	グラスプランス 440	うりくしょうとう
	TOTIVITUE		AGANTANAGG
	A C C C C C C C C C C C C C C C C C C C		TCTCCACCTT
	しじたるただらしても	17:511101	ACABARTACG
	1001	1011	

1321 GAAAGCTCCA TGAAAGCTGG TTGGGGACTA ATTTTGTAAC TACTTTATTC CCAGATCCTG CTTTCGAGGT ACTTTCGACC AACCCCTGAT TAAAACATTG ATGAAATAAG GGTCTAGGAC	1381 TAAITTCTCT AAATAAACCC TGGAATCTTG CCTTAICTCC TICAGGTTAA AAGCCAACTG
TACTTTATTC ATGAAATAAG	TTCAGGTTAA
atttgtaac Taaaacattg	CCTTATCTCC
TTGGGGACTA AACCCCTGAT	TGGAATCTTG
TGAAAGCTGG ACTTTCGACC	AAATAAACCC
GAAAGCTCCA CTTTCGAGGT	TAATTTCTCT
1321	1381

ATTAAAGAGA TITATTIGGG ACCITAGAAC GGAATAGAGG AAGTCCAATT TTCGGTTGAC	1441 CAAGGICTAA TGACTGCAGG ATCTAGCTAT CCATTGITIC TGGCCGCCTA TGCGTGCACT GTTCCAGAIT ACTGACGTCC TAGATCGAIA GGTAACAAAG ACCGGCGGAT ACGCACGTGA
AGTCCAATT	TGGCCGCCTA
GOAATAGAGG	CCATTGTTTC
ACCETAGAAC	ATCTAGCTAT TAGATCGATA
TTATTTGGG	TGACTGCAGG ACTGACGTCC
ATTAAGAGA TITATITIGG ACCITAGAAC GGAATAGAGG AAGA	CAAGGTCTAA GTTCCAGATT
1361	1441

1501 GGGTGTCTGG CAGAGAGCT GGGTAAATTG TAGTTTCATT GTAGCTGTCT GACTTGGATT CCCACAGACC GTCTCCCGA CCCATTTAAC ATCAAAGTAA CATCGACAGA CTGAACCTAA
GTAGCTGTCT
TACTITCATT
GGGTAAATTG CCCATTTAAC
CAGAGAGGCT GTCTCCCGA
GGGTGTCTGG
1901

1561 TCTCACGCCT ACTTCACTGG AAACGCAAAC TCTCACAGCA TTTTGTTTTA GTTTCAGAAT AGAGTGCGGA TGAAGTGACC TTTGCGTTTG AGAGTGTCGT AAAACAAAAT CAAAGTCTTA
AAAACAAAA
TCTCACAGCA AGAGTGTCGT
AAACGCAAAC TITGCGITIG
ACTTCACTOG TGAAGTGACC
TCTCACGCCT AGAGTGCGGA
1561

1621 CAGAGCAAAT TAGAAGTCTG AATTTCCTTC AACACTTGGA AATAATTTAT TIALLISMAA GTCTCGTTTA ATCTTCAGAC TTAAAGGAAG TTGTGAACCT TTATTAAATA AATAAACTTT
TTATTAAATA
AACACTTGGA TTGTGAACCT
AATTTCCTTC
TAGAAGTCTG
CAGAGCAAAT GTCTCGTTTA
1621

<sup>1681</sup> TATATTCATA ATTAATTCGT TATAAAATG TATTAAATGC TTATTTGAGT CAGCAGAGGA ATATTTTAC ATAATTTACG AATAAACTCA GTCGTCTCCT

FIG. 72D

TTCAGAACAT AACTCTTGTA TAGAGGAAAA ACGGAAGTAA TGCCTTCATT TAGAAGGTGG ATCTCCTTTT ATCTTCCACC TTTATGAAAG AAATACTTTC AGATAGAAAC TCTATCTTTG 1741

GATTATCTCA CTAATAGAGT AAAAGCAGGA TTTCGICCT TGTCATTTTA ACAGTAAAAT GAAACATTAA CTTTGTAATT CCCATTAGIT CTCOTTTACA GAGCAAATGT 1801

AAATATTTTG TITATAAAAC GTTGGATAAG TATCATTGAA CAGCAATACC CTTAGAATAA GAATCTTATT TAAAACATTT ATTTTGTANA 1861

AGTAGGCAAG TCATCCGTTC TTTCAGTGAA ALAGTCACTT CATGACTCTT GTACTGAGAA AATCTGTTTG TTAGACAAAC GCAACTTAAA GTTAACCANA CAATTGGITT

1921

AATTTGTGTT TTAAACACAA TAATATTGAT ATTATAACTA ATGTCAGAGG TACAGTCTCC ATCTCACCTA TAGAGTGGAT ATTCAGAAAT TAAGTCTTTA TCTTAATTT AGAAATTAAA 1981

TCGTATCTCA AGCATAGAGT ATCTATAGGC TAGATATCCG AATAAGTCCT TTATTCAGGA AATAATGAAA TTATTACTTT TACATACAAC ATGTATGTTG AATGITIATT TTACAAATAA 2041

2101 TGCCTATTTT TGGATGTATT TTTCA ACGGATAAAA ACCTACATAA AAAGT

## FIG. 73A

<b>9</b> —	TAT
	TATTTTT
0.	AGATAGGACT TCTATCCTGA
4	ACGASCCTAT
30	GCCATGAGAT CCGTACTCTA
50	TGAAAAATA' ATCAAAAATA GGCATGAGAT ACGAGCCTAT AGATAGGACT TATTTTTAT ACTTTTTATG TAGTTTTTAT CCGTACTCTA TGCTCGGATA TCTATCCTGA ATAAAAAATA
01	1 TGAAAAATAÓ ATCAAAAATÁ GGCATGAGAT ACGAGCCTAT AGATAGGACT TATTTTTAT ACTITTTATG TAGTTTTTAT CCGTACTCTA TGCTCGGATA TCTATCCTGA ATAAAAATA

ACATTAGGTG TGTAATCCAC AATTATCAAT ATTACCTCTG TTAATAGTTA TAATGGAGAC GIAMAACACA TGTATTATT ACATAATAAA TATTGTTGTA 61

ATGCAMACAG TACGTTTGTC AAAAAGAGTC TTTTTCTCAG ACTITICACTG TGAAAGTGAC AATITIAATT TCTCTTGCCT TTAAAATTAA AGAGAACGGA AGATATTCTG TCTATAAGAC 121

GATAGGTATT CAACTICAAT GTTGAAGTTA TITITITATC AAAAAATAG TTGCANANTA AACGTTTAI ATTITIANGT TGCAAACCAA TAAAAATTCA ACGTTTGGTT 181

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CCAGATGTTC GGGTGTCAAA TTCAACTAAT AAGTTGATTA CATTAATTGT GTAATTAACA CTAAGATATG GATTCTATAC GCTGTTAATT 241

ATCTTCCTCT TAGAAGGAGA GGCAAAAAGG AGATCCACCT TCTACTTTCA TAAAGTTTCT CCGTTTTTCC TCTAGGTGGA AGATGAAAGT ATTTCAAAGA TGAAAATGAA ACTTTTACTT 301

ATATTTCAMA TTAATACTTA TATAAAGTTT AATTATGAAT ATAACGAATT TATTGCTTAA TTATGTAMA AATACATTTT ATAAGCATTT TATTCGTAAA GCTGACTCAA CGACTGAGTT 361

CTGATTCTGA TAAGTCTGAA GATTAAACGA CTAATTIGCT GTTGAAGGAA ATTCAGACTT TANAGGTTCA CAACTTCCTT ATTTCCAAGT ATTTATTA TANATANAT 421

FIG. 73B

AAGTAGCGTG AGAAATCCAA TICATCGCAC ICTITAGGIT AATGCTCTGT GAGAGITTGC GTTTCCAGTG TTACGAGACA CTCTCAAACG CAAAGGTCAC 481 AACTAAAACA TTGATTTTGT

CAGACACCAG TGCACGATAG GTCTGTGGTC ACGTGCTATC GTCAGACAGC TACATGAAAC TACATTTACC AGCTCTCTGC CAGTCTGTCG ATGTACTTTG ATGTAAATGG TCGAGAGACG 541

AGACCTTGCA CTCAGTCATA GTAGCTAGAT CATCGATCTA CGCAGAACAT GCGTCTTGTA

601

AACCIGAAGG AGATAAGGCA AGATTCCAGG GTTTATTTAG AGAAATTACA TTGGACTTCC TCTATTCCGT TCTAAGGTCC CAAATAAATC TCTTTAATGT CAACCGAAAA CTTOOCTITY 199

GGATCIGGGA ATAAAGTAGT TACAAAATTA GTCCCCAACC AGCTTTCATG GAGCTTTCAA CCTAGACCCT TATTTCATCA ATGTTTTAAT CAGGGGTTGG TCGAAAGTAC CTCGAAAGTT 721

TTCTAGTICT TAATCGCATG CATACAATGC ACATACATAT ATACATGCATA AAGATCAAGA ATTAGGGTAC GTATGTTACG TGTATGTATA TATGTACGTA TTATTAATTA AATAATTAAT 781

ACCIDIDGAT AAAACAGAAA TGGACACGIA TITIGICITI ATGAITGGAC GCAAACGGAA ATAAGAITCC TACTAACCTG CGITTGCCTT TATTCTAAGG TAATTTTATG ATTANAMIAC 178

NNNNNNNNN NNNNNNNNN GACGAGATON CTCCTCTACN CCACACTGAG GGTGTGACTC GAGTGAGGGA TCAGGAAACA CTCACTCCCT AGTCCTTTGT CTGAACCAAT GACTIGGITIA 106

NTAGTGGGTG GGGGGGGGAC ATCAATAAAG AACTCTTCTG TGTCAGCCAC TGAGCACGGA NATCACCCAC CCCCGGCCTG TAGTTATTC TTGAGAAGAC ACAGTCGGTG ACTCGTGCCT 196

CAGATGAAGA CICIACITCI ATAAAGGGAT GAGAGTGAGG GCAANTACCA GAAGAATAAA ATCCTTTTAA TATTTCCCTA CTCTCACTCC CGTTNATGGT CTTCTTATTT TAGGAAATT 1021

ATCTITIMAC AACCCCAAGG TOAAGCTAGT TAGAAAATTG TIGGGGTTCC ACTICGAICA CACAGTGTGT GGNTTCAAAA GTGTCACACA CCNAAGTTTT GTGTCACACA TTGTTATGAG AACAATACTC 1081

TOGRAGATAT TTGAATTTGT TTAAACCCAT CTGGTCCTAG CCCTATTCTT TGAATCCCGA ACCTTCTATA AACTTAAACA AATTTGGGTA GACCAGGATC GGGATAAGAA ACTTAGGGCT 1141 TGGAAGATAT

## FIG. 73D

ACTAGTCCTG TGATCAGGAC GCADDADTO ACTACCTGOT GATACCTTAG COTCCTCACC TGATGGACCA CTATGGAATC AGAATTCCGA TCTTAAGGCT AAGAGGGTCA TTCTCCCAGT 1201

CGAAAATCCC AGTATCITGG TAAAATAATA AATAAAGTCC TCATAGAACC ATITTATTA TTATTTCAGG TOTATTAKAG TCCAATGAGG ACATAATTTC AGGTTACTCC TOTATTANAG 1261

AATTTGCAGA TUNNNINNT AATTIGCAGA AHNNNINNNA TIAAACGICT ACATGCIATA TIATITACIA TGTACGATAT AATAXATGAT AGTACTGTGC TAGGAGATTI TCATGACACG ATCCTCTAAA AGTACTGTGC 1321

CATTGAACAA GIAACTISIT GAGGGACTCO CTAACGCTGA AATAGGGTAA TTATCCCATT CTCATCATA GAGTAGTATT TAATATTATC ATTATAATAG 1381

AAIAAAAGAG TCTAGCTTGC TTATTTTCTC AGATCGAACG GACCTTAMA CTOGAATTTT AAAGTCAAAA TTTCAGTTTT AAGAAGTGGC TTCTTCACCG CAAGGCCACT GTTCCGGTGA 1441

CCCAGGAAAA CANATCAGTA GTHTAGTCAT Gannaagtet Ctnnttcaga CTGCTTTTCT TAGAAAGTTG GACGAAAAGA ATCTTTCAAC GACACACCAA CTGTGTGGTT 1501

1551 ACAGCAAAAG ACCCGCTGGT AAAGACCTGT CCAGATTGCT GACCTGGTTC ACACANITCC

FIG. 73E

TGTCGTTTTC TGGGCGACCA TTTCTGGACA GGTCTAACGA CTGGACCAAG TGTGTNNAGG

AAAACAAACA TTTTGTTTGT GAGAGGTAAA AAGCITGCCT CTGTTACITC CAA3GAAGAA AGAATGCACA TTCGAACGGA GACAATGAAG GTTCCTTCTT TCTTACGTGT AAGCTTGCCT 1621

AAACTTCCTC AAGCAAAAA TTCGTTTTTT AACCAAACAA AACAAAACAA AACAAAACAA AACAAAACAA TIGGTTIGIT TIGITIIGIT TIGITIIGIT

1681

CTIGGAACCT ICCTACGICC TANITICAGG ITCICICAGI GAACCTIGGA AGGAIGCAGG AINAAAGICC AAGAGAGICA TGTCTTGCAG GGCTCCAGCA ACAGAACGTC CCGAGGTCGT 1741

CTGTCCTACC AGCAGCTTGT CGAGAACTCA GCCCTGCACC GACAGGATGG TCGTCGAACA GCTCTTGAST CGGGACGTGG ACCTGAGTGA TGGACTCACT TCTACCCTCA 1801

1861 GTTCCCAGCT ACCCTCCTCC TAACTCGASG GGTGCT CAAGGGTCGA TGGGAGG ATTGAGCTCC CCACGA

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9	AGACTCAT
<b>6</b>	GGATICIGIT GAGCCTAGE TCATTATGAT GICCIGTIGT CCIACCCAAA TAAGACTAT CCIAAGACAA CTCGGGATCG AGTAATACTA CAGGACAACA GGATGGGTTT AITCIGAGIA
<b>-</b>	GTCCTGTTGT CAGGACAACA
0-	TCATTATGAT AGTAATACTA
20	GAGCCCTAGC CTCGGGATCG
10	1 GGATICTGIT CCIAAGACAA

ATAXATAAT TATTTATTTA TALALALAN CCCAACTACA TCICAATAAT TAATGAAGAT GGAAATGAGG GGGTTGAAGT AGAGITATAA ATTACTTCTA CCTTTACTCC 61

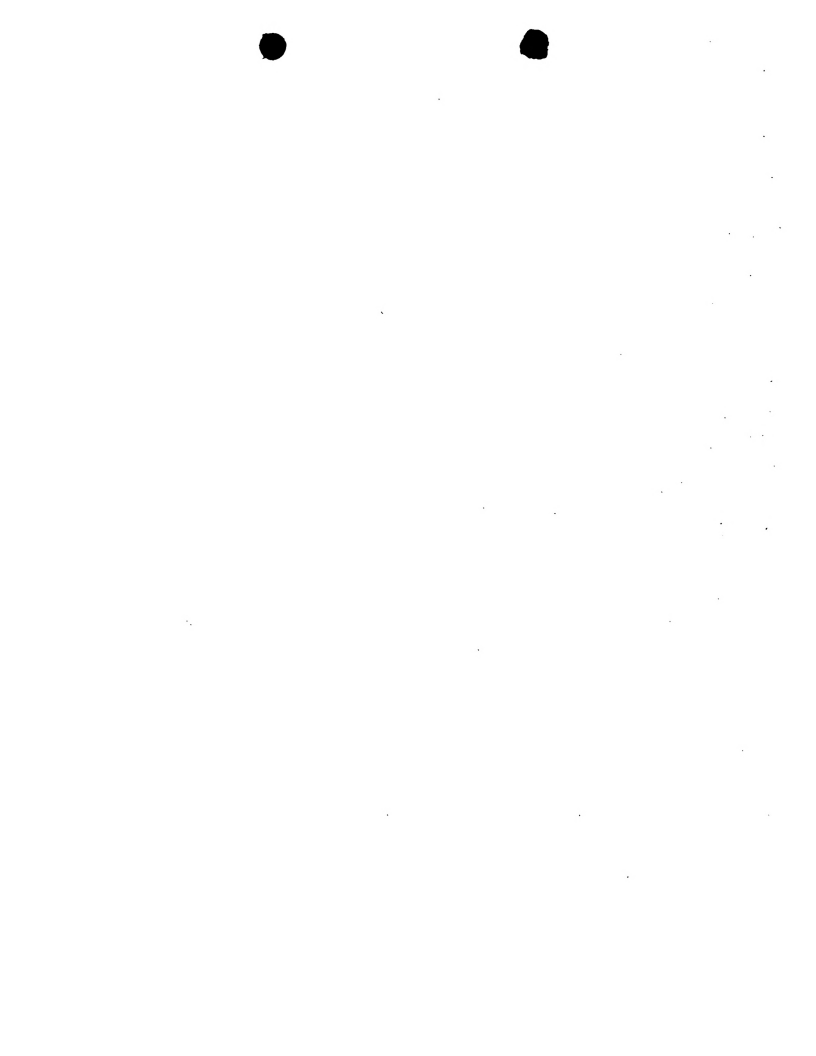
CITCIAIGAA ATAATUTICI GAAGATACIT TATTACAAGA TTTCAAATAC AAAGTTTATG TTCCCCCCA TTTATTATTT AAGGGGGGT AAATAATAAA AAAAGAAACA TTTTCTTTTGT 121

CTTTANTATC GAATTTATAG CTGTGAATAC ANATHITAM AGNANICAM ATTAITIGGAN TITAINAMIN TCIFTAGITA TANIMACCIT TAGGGAGAGA ATCCCTCTCT 181

GAAGTCGGGA ACTEGETTTA TGACCCAAAT CTTTCCTATO ATGTTGAGTT GAAAGGATAC TACAACTCAA TCATTATCCG GTGTCAACTA AGTAATAGGC CACAGTTGAT 241

CACACCAATA TCAAATATGA TATACTTGTA GTGTGGTTAT AGTTTATACT ATATGAACAT AATAATGCTG TAAANNNNNN AGTTAGTCTA TTATTACOAC ATTINNNNN TCAATCAGAT 101

CHARCETTAT ANAGREGET CTTITITICE TITITITITE CTATOLANAIA TITICICCAL GALANALAGA ALANAMAAA AACCTCCAAG CATAAAAAAA TTGGAGGTTC GTATTTTTTCT 161



## FIG. 748

481 TGCAACCICC ACCICCCAIG ITCAAGGAT TCTCCTTCCT CAGICTCCTG AGTAGCTGGG	ACCTIVIDADO TODAGOTAC AAGTECCETA AGAGGAAGGA OTCAGAGGAC TCATCGACCC
CAGTCTCCTG	<b>OTCAGAGGAC</b>
TCTCCTTCCT	ACACCAACCA
ITCAAGGGAT	AAGITCCCTA
ACCICCCAIG	TOGAGGGTAC
TOCANCCTCC	ACCTIOGAGO
481	

541 ATTACAGGTG TGCACCACCA CACCCAGCTA ATTITITATA TTTTATAGA GACAGGGTTT	TAATGTCCAC ACGTGGTGGT GTGGGTCGAT TAAAAACATA AAAATTATCT CTGTCCCAAA
TTTTAATAGA	A.A.T.TATCT
ATTTTTCTAT	TAMMACATA
CACCCAGCT'A	GTGGGTCGAT
TOCACCACCA	Acordorage
ATTACAGGTG	TAATGTCCAC
541	

CCCGCCTCAG	GGGCGGAGTC
601 CATCGATGIT GGCCAGGCTA GTCTCGAACT CCTGACCTCT AGGIGATCCA CCCGCCTCA(	GTAGCTACAA CCGGTCCGAT CAGAGCTTGA GGACTGGAGA TCCACTAGGT GGGCGGAGTC
CCTGACCICT	GGACTGGAGA
GTCTCOAACT	CAGAGCTTGA
GGCCAGGCTA	CCGGTCCGAT
CATCGATGTT	CTAGCTACAA
601	

661 CCTCCCAAAG TIGTAGAAIT ACACGIGIGA GGCACTGCTC TGGCCAGGAG ATACATITITI GGAGGGTTTC AACATCTTAA TGTGCACACT CCGTGACGAG ACCGGICCTC TAIGTAAAAA

721 GATAGGITTA ATTIATANA ACACTGCACA GATTIGGACT TGCTGGGNAA TCACGATCCA CINTCCAAAT TAAATATTIC TGTGACGTGT CTAAACCTCA ACGACCCTTT AGTGCTAGGT

# FIG. 74C

99	ပ္ပ
ATTGATCA	TAACTAGI
T'INTAICTCA	AATATATAGT
TACITAATSA	ATGAATTACT
TTTTTATTGG	MANATANCO
GACCCAGCAA	CTGGGTCGTT
781 GTATGCATTT GACCCAGCAA TTTTTATTGG TACITAATSA TTATATCTCA ATTGATCAGG	CATACGIAAA CIGGGICGII AAAAATAACG AIGAATIACT AAIAIAGAGI INACTAGICC
781	

GAGGCAAGGT	CICCGITCCA
<b>OGACAGITITO</b>	CCIGICAAAC
ACATITGAGA	TCTAAACTCT
TTGTGTGTGG	AACACACACC
TGCGAAGAAT	ACCCTTCTTA
841 TIGAACTCTG TGCGAAGAAT TTGTGTGTGG ACATTTGAGA GGACAGTTTG GAGGCAAGGT	AACTTGAGAC ACGCTTCTTA AACACACCC TGTAAACTCT CCTGTCAAAC CTCCGTTCCA
841	

ACTGAGAAAS	TCACTCTTTC
TOGGSCATAT	ACCCCCTATA
<b>OTTTOCAAGT</b>	CANACOTTCA
TTTCAATCTT	AAACTTAGAA
ATTTANAGAA	TAMPLITCIT
901 ATITIAGIAG ATITAAAGAA TITCAATCIT GITTGCAAGI TGGGGCATAT ACTGAGAAAS	TAMANICAIC TAMAITICIT AMACITAGAA CAMACGITCA ACCCCCIATA TGACTCTITC
106	

IÇ Y	Nor
TCAMON	ACTITICE
ATCHTTATA	TACAACTTAT
ATTATOATOT	TANTACTACA
TCATATATT	ACTATATANA
GCAGATAAAT	CGTCTATTTA
961 AGAAGACAAT GCAGATAAAT TGATATATTT ATTATGATGT ATGTTCAATA TGAAAGATCA	TCTTCTGTTA CGTCTATTIA ACTAINTAAA TAATACTACA TACAAGTTAT ACTTTCTAGT
961	

1021 CAAATATAA CATACATNNA TCTTACTTAA CATACCTCAG TTTTAGAGCT ACCGTATGTA	GITITAIAIT STAIGIANNI AGAATQAAIT GIAIGGAGIC AAAAICTSGA IGGCATACAI
ITTIAGRECT	AMATCTCGA
CATACCTCAG	GTATGGAGIC
TCTTACTTAA	AGAATOAATT
CATACATNNA	STATGTANNT
CAMMATATA	GITITAIATT
1021	

GCACTCTTAA	CITCICAGGI AAAGATAAAI CCAIICAAGG AAATCAGGAA AAIAAIGACC CGIGAGAATI
TEATTACTOR	ATA TCACC
FITAGICCT	AAATCAGGAA
GGTAAGTTCC	CCATTCAAGG
TTTCTALTTA	AAAGATAAAT
1081 GAAGAGICCA ITTUTATITA GGIAAGIICC ILIAGICCII TIATIACIGG GCACICITAA	CTTCTCAGGT
1081	

1141 TTACATGING CITGRAATAT GICCAGIITG AGCAGIGAAC IGAAAATGIC ATGIGATTAA	ANTOTACATO GANCTITATA CAGGICAAAC ICGICACTIG ACTITILGAG IACACTAATI
TGAMAATGTC	ACTITIZACAG
AGCAGTGAAC	TCGTCACTTG
GICCAGITIC	CAGGTCAAAC
CTTGAAATAT	GAACTTATA
TTACATGTAG	ANTOTACATO
1141	)

<sup>1201</sup> GTACATATAT ANTITITIT CATAGIAGGT CAATAACCTC CITITATIGA CTAATGAATC CATGIATATA TIAAAAAAA GTAICAICCA GTIATIGGAG GAAAAIAACT GAITACITAG

<sup>1261</sup> ACTICICIA TGATTATAGG TCAAGAGATT ACTAATATGC

## FIG. 75A

0,	AATGAATATT TTACTTATAA	TATCAGATA	GATTCTGTC FCTAAGACAG	ITAACTAAAA FATTGATTTT
<b>8</b> 0 -	1 aatcaaaata aaacagttaa agtttgatta ctataatcaa acacaaaaa aatgaatatt	61 ATCITITATG TCAGTAGAGG GTGAATGAAT CCITCAGGAT ITTGATGATA GTATCAGATA	121 CCCAGCACTA TGCTAGAAGT TGTGAAGAAT TCACGAGATG AATAAATCAC AGATTCTGTC	181 CTCAAAATGG TTAGATCTAT TCAGGAAACA AAGCTAAAAA AACCCCACCA ATAACTAAAA
	Ttagttttat tttgtcaatt tcaaactaat gatattagtt tgtgtttttt ttacttataa	TAGAAAATAC AGTCATCTCC CACTTACTIA GGAAGTCCTA AAACTACTAT CATAGTCTAT	GGGTCGTGAT ACGATCTTCA ACACTTCTTA AGTGCTCTAC TTATTTAGTG TCTAAGACAG	GAGTTTTACC AATCTAGATA AGTCCTTTGT TTCGATTTTT TTGGGGTGGT TATTGATTTT
40	CTATAATCAA	CCTTCAGGAT	TCACGAGATG	AAGCTAAAAA
	GATATTAGTT	GGAAGTCCTA	AGTGCTCTAC	TTCGATTTTT
30	AGTTTGATTA	GTGAATGAAT	tgtgaagaat	TCAGGAAACA
	TCAAACTAAT	CACTTACTTA	Acacttctta	Agtcctttgt
20	AAACAGTTAA	TCAGTAGAGG	TGCTAGAAGT	TTAGATCTAT
	TTTGTCAATT	AGTCATCTCC	ACGATCTTCA	AATCTAGATA
10	aatcaaaata	Atchithig	CCCAGCACTA	CTCAAAATGG
	Ttagttttat	Tagaaaatac	GGGTCGTGAT	GAGTTTTACC
	ત	61	121	181

115/130

301 AGAGGAGGIA AAAAGAIAAC TCTTCCAAAA GGAATACTAT AIACTGIAAA CTGTGTACTG

241 ATCAACCAAA TGAAAAACAA CAATCATAAA ATAAGTAAGT ACCTATAGAA AGAAAAGCTC TAGTTGGTTT ACTTTTGTT GTTAGTATTT TATTCATTCA TGGATATCTT TCTTTTCGAG

# FIG. 75B

CTCATCAATT	GAGTACTTAA
421 CACAAGCCTA AATATGTAGT TGCTTCACAG AAGGTTAGAA GTAAATTAAC CTCATGAATT	GIGIICGGAI TIAIACAICA ACGAAGIGIC TICCAAICII CAITIAATIG GAGIACIIAA
AAGGTTAGAA	TICCAATCIT
TGCTTCACAG	ACGAAGTGTC
AATATGTAGT	TTATACATCA
CACAGCCTA	GTGTTCGGAT
121	

GAAAGATTIT AATACCAAAT CTTTCTAAAA ITAIGGITTA TCTTGAGAGA ACTTGTAAGG ACTAAGCTTT CGATTTTGGA AGAACTCTCT TGAACATTCC TGATTCGAAA GCTAAAACCT 481

TGCTTAGATA ATACCTAGGA ACGAATCTAT TATGGATCCT AATCTCAATC ATTATAATAG TTAGAGTTAG TAATATTATC AAAAAGTACC TTTGTTTGGT TTTTTCATGG AAACAAACCA 541

116/130

ACAMATTAMA TATTAMATTT ACTTTAMAMA ARAGTACATG ATTGGGGAAT CACAACTGGC TOTTTAMTTT ATAMTTTAMA TGAMATTTTT TTTCATGIAC TAMCCCCTTA GTGTTGACGG 601

AAAAACACTG AACCAAATAT TTTTTGTGAC TTGGTTTATA

GAAAAGAATG

NATATGCACT

GAATGATCTA AGAGANNNN NTATACGTGA

TCTCTNNNN

CTTACTAGAT

199

721

TTAAATTGGA AAAAAATAGT AAGGAATATC AGAAGCAAAA AATTTAACCT TTTTTTATCA TTCCTTATAG TCTTCGTTTT AAGTTTAAAA TTCAAATTTT NTGTTTTTT NACAAAAAA

CTTAGATGGA TCCTCAGAGG TAGCACGAAA TTTGGCTTTG AGGAGTCTCC ATCGTGCTTT AAACCGAAAC AAAGCAAGAA TTTCGTTCTT TITATITAC AAATAAAATG 781

FIG. 75C

GGTTCACATA GTTTAAAGCT TCAGGAGTTA TGAAAAGGAT ACTTTTCCTA CTATGGCCCA TCTATCAAAGAGAGAGAGAGAGAGAGAGATAGTTTC 841

TCCTGACCAG ACAACAATA TTGTTGTTAT GTGGTCTANG CACCAGATTC GTGCATAAAG CACGTATTTC GCAGAAGACT TACCTTAGAT ATCGAATCTA 106

GATCACGAGG AAGGIGGGIG TCACNCTNAA TNCCAGCACT TTGGGAGCCC AGTGNGANTT ANGGTCGTGA AACCCTCGGG CACTCCCCG GTGAGGGGGC

961

AAAAATAGAA TTTTTATCTT CCTCTCTACT GGTGAAACCG TGACCAACAT QAGACCAGCC CTCTGGTCGG TCAGGAGTTT 1021

AGACAGGAGA TCTGTCCTCT CAGGAGACTG CAGCTGAACT CTTCTAATCC NGCCTACGTG NCGGATGCAC AATTAGCCG TTAATCGGC 1081

ACTCCAGCCT AGGGTGCAAA TGAGGTCGGA TCCCACGTTT NNGCCACTGC NNCGGTGACG AAGCTTNNNN TTCGAANNN CCCAGCATGC GGGTCGTACG ATCACTTGAA 1141

1201 AAAAAAAA ANGACACATT ACTCAGGTAA GGTAATCAAT AA TITITITITIT INCTGTGTAA TGAGTCCATT CCATTAGITA TT

### FIG. 76A

-	AAGG	ŢŲ	AAA	TTAT	111	111	111	TCT	CCCC	CAA	rgta		GIT	 ATAG	-
-	T000       T000	III	III					III		111		IIII	III		-
-	ATTT	ĬĬĬ	111	1111			IIII			Ш		IIII	III	1111	-
-	TAGG														-
<u>-</u> -		TH	111	$\Pi\Pi$	$\Pi\Pi$			ĪĬĬ	1111	111	1111		III		-
-	1111	HII	HI		CATO	1:11	IIII	ĪĪĪ		111		HIII		CTTT	-
-	IIII		111			111	HII	ĪШ	1111	111	1111	ĪШ	$H\bar{\Pi}$	ATGA      ATGA	-
-	ATTO		$\mathbf{H}\mathbf{H}$	1111		1111	1111	111		$\Pi\Pi$	1111	$\Pi\Pi$			-
-	TTAC	111	111	1111		$\Pi\Pi$	1111	111	HH	111	1111	$\mathbf{H}\mathbf{H}$		TTAA TTAA	<u>-</u>
_	3.000	WII 2 2		3 707	-	-	>~~	720		120	1001	3.44	MCC		_

### FIG. 76B

-	ATT	FAAAA	AATT	CCCTI	TCGA	TGTA	GAACA	AATAG	388777	GCCI	) 3T -
-	GGGG'	CTAC             CTAC	TTGC      TTGC	TTATI       TTATI	ATAT      ATAT	PTGTA        TGTA	AGCTA       AGCTA	CTGCT)       CTGGT)	AGGAAA         AGGAAA	TAGCA       TAGCA	W -
										GAGGA!       GAGGA!	
		$\mathbf{H}\mathbf{H}$	1111	11111			11111			AGATG!        AGATG!	ΙĪ
		1111	1111		HH		11111		111111	GGTAC:	ΙĬ
										AACCA       AACCA	
	TTAN	11111	1111	11111	1111	HHH	11111	1			

# FIG. 77A

9-	GTAATATC
50	AGANAACACÁ GTGTCTITCT TTCCTTATTT TAANTTGGTT GTTCCAGATT CGGTAATATÓ TCTTTTGTGT CACAGAAGA AAGGAATAAA ATTTAACCAA CAAGGTCTAA GCCAFTATAG
<b>4</b>	TAMATTGGTT ATTTAMCCAN
30	TTCCTTATTT AAGGAATAAA
20	GTGTCTTTCT CACAGAAAGA
0.	1 AGAAAACACÁ TCTTTTGTGT

TTCTCATTAG	IAA AAGAGTAATC
CTTCAACCTT	GAAGTTGGAA
AGAACTITAT	TCTTGAAATA
AATGAGTACC	TTACTCATGG
ATTACACTTA	TAATGTGAAT
I AATITICAAT AITACACITA AATGAGTACC AGAACTITAT CITCAACCIT ITCICATTAG	TTAAAAGTTA TAATGTGAAT TTACTCATGG TCTTGAAATA GAAGTTGGAA AAGAGTAATC

121 GCCTACAACA AAGGACATCT CGGATAGAAT TTCCCTTTTC TTTTTGCTAC TATAAGCTCT	CGGATGITGI ITCCIGIAGA GCCIAICITA AAGGOAAAAG AAAAACGAIG ATAITCGAGA
TTTTTGCTAC	AAAAACGATG
TTCCCTTTTC	AAGGGAAAAG
CGGATAGAAT	GCCTATCTTA
AGGACATCT	TTCCTGTAGA
GCCTACAACA	CGGATGTTGT
121	

181 AAAAATCCTC AGAACATCAG ATITAGAAAT GTTCTIAITA GTGGTAGTGA GCATTTGCTA TTTTTAGGAG TCTTGTAGTC TAAATCTTTA CAAGAATAAT CACCATCACT CGTAAACGAT
GTGGTAGTGA
GTTCTTATTA
ATTTAGAAAT TAAATCTTTA
AGAACATCAG TCTTGTAGTC
AAAAATCCTC TTTTTAGGAG
181

AAATTCCTAT	TTTAAGGATA
IC CCCACAGGCC AAATTCO	GGGTGTCCGG
A GCANGTAGAC	T CGTTCATCTG
AATATAATAA	TTATATATT
CA CTAGCTTACA AATATAATAA GCAAGTAGAC	CATCGAATGT
41 TTTCCTACCA	AAAGGATGGT GATCGAATGT TTATATTATT CGTTCATCTG GGGTGTCCGG TTTAAC
241	

AGAMAATAT	AACAAGATGT CAGCTTTCCC TTAAAAATT TTAAATTAAA
CCCACTAAAG	GGGTGATITC
AATTTAATTT	TTAAATTAAA
AATTTTTAA	TTAAAAATT
GTCGANAGGG	CAGCTTTCCC
01 TIGITCIACA GICGAAAGGG AATITITIAA AATITIAATIT CCCACTAAAG AGAAAAIAI	AACAAGATGT
301	

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TTTCCCTCA1	AAGGGAGTA
TGTAAATTGT	ACATTTAACA
ATTTGCTATG	TANACGATAC
TAATTTTAA	ATTAAAAATT
CANATGACAG	PANTIGITIA GITTACIGIC ATTAAAAIT TAAACGATAC ACATTTAACA AAAGGGAGIA
361 ATTAACAAAT CAAATGACAG TAATTTTTAA ATTTGCTATG TGTAAATTGT TTTCCCTCAT	TAATTGTTTA
361	

<sup>421</sup> TATTTATAAC AATTCATACT ACAAITTAAT TTAGTAAACA TTTTTGTAGA AAATATTTAA ATAAATTT AAAAATATT TTATAAATT

FIG. 77B

CAGTGCATGC TTCTTGTAGG CCACAGCCAT GTCACGTACG AAGAACATCC GGTGTCGGTA CAGTGCATGC AACAAAGATA CIGAAAGITA ATATNAAACC TIGITICIAT GACTITICAAT TATANTITGG 481

ATCTACACIG GCCAAATTCC TAGATGTGAC CGGTTTAAGG CACAGAAAA TITGITCTGT TACTCTAAAC GTGTCTTTTT AAACAAGACA ATGAGAITTG AACCTGTAAG TTGGACATTC

541

601

GTCCTCTCTG TAAGGTGGGC CAGGAGAGAC ATTCCACCCG TTTAACCCCG GGATATAACC TAGTAAATGT AAATTGGGGC CCTATATTGG ATCATTTACA AATGCTCGAA TTACGAGCTT

GATTCTACAC CTAAGATGTG TTAAGAMAT AATTCTTTA ATACAAGAAA ATAATGGTAT TCATAAAGTT TATGTTCTTT TATTACCATA AGTATTTCAA ATGTCACAGA TACAGTGTCT 661

AAAAAGAGAT AATTTTTACC TTTTTCTCTA TTAAAATGG CACTATAACT TITTACATTG GGGAGAGAA GTGATATTGA AAAATGIAAC CCCCTCTCTT ATGTAAAACC TACATTTTGG 721

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9	Atctccattt Tagaggtaaa	titatcagta taaaatagaa aaatagecat attitatcet	CAGCCATGAA GTCGGTACTT	CCAGACATTG GGTCTGTAAC	CAAGGATACT GTTCCTATGA	GCCAGGGACT
0£0	TTAGGCTTTT AATCCGAAAA	tttatcagta <b>aaa</b> tagtcat	TAGTGTATGT ATCACATACA	AGTCATTGCT TCAGTAACGA	CTCAGTGTGG GAGTCACACC	GCAAACTTTG GCCAGGGACT
40	TTTTGAAATG AAAACTTTAC	TCCTTCTTAT ACGAACAATA	CATGAGTATC GTACTCATAG	AACTICAGGG AACCTAATIG TTGAAGTCCC TIGGATTAAC	CGGGCAATGA GCCCGTTACT	CTCCTCTGAT
30	TTATTGACAG AATAACTGTC	AAATTITCCA ACATGGGTGT TGCTTGTTAT TITAAAAGGI TGTACCCACA ACGAACAATA	TTAGTATATA AATCATATAT	<b>aa</b> cttcaggg Ttgaagtcc	TNNNNNNCT	TGTTTCTGGA AGGCACTGGA ACAAAGACCT TCCGTGACCT
50	GGGCAATTTC		GTTCTGGAAT Caagacctta	TCAGATGTTT AGTCTACAAA	CCCACTATAT GGGTGATATA	
10	GATGCTATT CTACGATAAA	TTTAGTACTT AAATCATGAA	121 GAGTGGTTCT CTCACCAAGA	AATGAACCTT TTACTTGGAA	TTGCTTTGAA AACGAAACTT	ACTGCAGGCC TGACGTCCGG
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421 TATTCAATAT TAGACTACAA GCAGTCTAAG GACTTCTCAG GGTTTCTAGC TCTCTCAT	ATAAGTTATA ATCTGATGTT CGTCAGATTC CTGAAGAGTC CCAAAGATCG AGAGAGATA
GGTTTCTAGC	CCAAAGATCG
GACTTCTCAG	CTSAAGAGTC
GCAGTCTAAG	CGTCAGATTC
TAGACTACAA	ATCTGATGTT
TATTCAATAT	ATAAGTTATA
421	

TGGGGCCAGA	Accedence
481 IICACACAIG CIIICCIAGI AAICICIACI CAIAICII ACIGCIACGC IGGGGCCADA	AAGTGTGTAC GAAAGGATCA TTAGAGATGA GTATATAGAA TGACGATGCG ACCCCGGTCT
CAIATATCTT	GTATATAGAA
ATCTCTACT	TTAGAGATGA
CITICCIAGI	GANGGATCA
TICACACATG	AAGTGTGTAC
195	

541 TAACNNNNNN CTTCCATTTT GTTTTTATCT CTATTCTTCT TCCCCTTCTG CTTTCATTAT	ATTCNNNNN GAAGGTAAAA CAAAATAGA GATAAGAAGA AGGGGAAGAC GAAAGTAATA
recentera	AGGGGAAGAC
CTATTCTTCT	GATAAGAAGA
GITTITAICT	CAAAAATAGA
CTTCCATTT	GAAGGTAAAA
TAACNNNNNN	ATTGNNNNN
541	

ACCTGGCATT	TGGACCGTAA
GTTCTGCTTA	CAAGACGAAT
TCCCAGATTT	AGGGTCTAAA
ATTGANACTT	TAACTTTGAA
TGCTTTCATT	ACGANAGTAA
601 IGAAACTITC IGCITICATI AITGAAACTI ICCCAGAITI GIICIGCIIA ACCIGGCAII	ACTITGAAAG ACGAAAGTAA TAACTITGAA AGGGTCTAAA CAAGACGAAT TGGACCGTAA
601	

TII	*
TTTTTT	AAAAAA
CATGICCTIT ITTITITI	GTACAGGAAA
CICCCATTGC	GAGGGTAACG G
GTGCTGCTTT	A GOAGAAGGGA CACGACGAAA GAGGGTAACG GTACAGGAAA AAAAAAAA
CCTCTTCCCT	GGAGAAGGGA
661 GGAACTGTTT CCTCTTCCCT GTGCTGCTTT CTCCCATTGC CATGTCCTT	CCTTGACAAA
661	

<sup>721</sup> ITITITITIT TOAGACAGIG TCACTCTGTT GCCCAGGCTG GAGTGCAATG GTGCCATCTT AAAAAAAA ACTCTGTCAC AGTGAGACAA CGGGTCCGAC CTCACGTTAC CACGTTAGAA

FIG. 78C

781

AGTAGAGATH TCATCTCTAN TTGTATTTTT CCGCTCATTT CCACTATOCC GGTGATACGG CAGGIGCCCA GCTGGGATTA

841

106

GTGANTCCGC CACTNAGGCO CCTSACCGCA GTCTCGAACT CTAGTCCGAC GATCAGGCTG GTGGTANCGA CACCATNGCT NNNNNNTTT CAGGCATGAG TCACTGCGNC CAGCCACCAT GTCCGTACTA GTCCGTACTA GCTGACATTA CCTCCTTGGC CTCCCAAGT GGAGGAACCG GAGGGTTTCA 196

INTICTCING AGGIGAGA ACACTGGCTC TICTAACAAG TIGAAATTTG ATAGAGACC ATAAGAGATC TCCACTCTCT TGTGACCGAG AAGATTGTTC AACTTTAAAC TATCTCTGG 1021

### **A**6

9		ATGTTAATGG	TACAATTACC
000		1 CACAAAAAA GATTATTAGC CACAAAAAA CCTTGAAGTA ACGCATTAAA ATGTTAATGG	GIGITITITI CTAATAATCG GIGITITITI GGAACTICAI IGCSTAATIT IACAATTACC
•	_	CCTTGAAGTA	GGAACTICAT
00		CACAAAAAAA	GIGTTTTTT
30	_	GATTATTAGC	CTAATAATCG
10		CACARADAA	GICTITIFIE
		-	

TTA TIDARCATCT GCTCATANTA CTITAATDAG TGCAAAGTGC TTTGAATATA	MACTEATAT
TGCAAAGTGC	ACGITTCACG
CTTTAATGAG	GANATTACTC
<b>GCTCATANTA</b>	CGAGTATTAT
TTGAGCATCT	AACTCGTAGA
61 ATTCACTTTA	TAAGTGAAAT AACTCGTAGA CGAGTATTAT GAAATTACTC ACGTTTCACG AAACTTATAT
-	

121 ATACGTCATT TAAACCTTAC CATAATICI'G AGGAATIGCT ACCTCCACTT CACAGATGGG	TATGCAGTAR ATTTGGAATG GTATTAAGAC TCCTTAACGA TGGAGGTGAA GTGTCTACCC	
ACCTCCACTT	TCGAGGTGAA	
ACCANTICCT	TCCTTAACGA	
CATAATIICIIG	GIATTAAGAC	
TAMCCTTAC	ATTTGGAATG	
ATACGICATI	TATGCAGTAA	
121		

181 GCACAGGAGG CTTAGATAAC ATGCCCAAAG TCATGCTTCT AGTAAATGGA TATAATTAAG CGTGTCCTCC GAATCTATTG TACGGGTTFC AGTACGAAGA TCATTTACCT ATATTAATTC	
AGTAAATGGA TCATTTACCT	
TCATGCTTCT AGTACGAAGA	
atgece <i>kaa</i> g taegggttpe	
CTTAGATAAC GAATCTATTG	
GCACAGGAGG CGTGTCCTCC	
181	

301 CGCTTTCCAG AGCATGTGCT GTTGATAGAG CTTGATGTCT AACTGTGTA AATTTTCCAT	GCGAAAGGIC ICGIACACGA CAACIAICIC GAACIACAGA TIGAGAGGIC TIAAAAGGIA
CTTSATGICT AAC	DAACTACADA TEG
GTTGATAGAG	CAACTATCTC
AGCATGTGCT	TCGTACACGA
CGCTTTCCAG	GCGAAAGGTC
301	)  - 

<sup>361</sup> TCTIATITGT CICACTGGTA TATAGITATT TITTACTACT TICATACACC TACTAAGAAG AGAATAAACA GAGIGACCAT ATATCATAA AAATGATGA AAGTATGTGG ATGATCTTC

FIG. 79B

AGCTTCACGT ATTITAATTC TCGAAGTGCA TAAAATTAAG GAATGCCTAA ATTTCATTTA TAAAGTAAAT 421 ACAGGAGGAT CAAAGATAGG TGTCCTCCTA GTTTCTATCC

CCTGGTTATC TITCAGCAGG GGACCAATA AAAGTCGTCC CACCAGIATA IGCCATGGIC GIGGICATAI ACGGIACCAG AGAATAAGAT ICAGGCAGAC TCTIAITCIA AGICCGICIG 401

TOACCOAGAA ADAMACATG GTAATGIITA TGAAATGGTG GGTICTTGTA GTITCACTTC ACTGGCTCTT TCTTTTGTAC CATTACAAAT ACTTTACAC CCAAGAACAT CAAAGTGAAG 111

TATTCTTGAT ATGGGCATGT ATAAGAACTA TACCCGTACA CCTTTACTGF ATTAGGATGA TGGATTAACT GGALATGACA TAATTCTACT ACCTAATTGA AACATATCTG TTGTATAGAC 601

TGTGTTTCCA GACAACTTA ACACAAGGT CTGTTTGAAT GAGAGACAAA ANACATAT ACTITIACTA AACAGCTACA TITIGITATA TGAAAATGAT TIGICGATGT 661

AGAGACIGAG IGIICAAACI GAATAATUTC GACCITAATI GIAACIAIAI TITAIGAAAT TCICIGACIC ACAAGIITGA CITATTAGAG CIGGAATTAA CATIGATAIA AAAIACITIA 721

CCTGTTAN	AGGACAATN
2A GACTICITIG GGCCIACCAC GGGCAITITG ITCCIGITA	GGTCGACATT CCGTTTTTGT CTGAAGAAAC CCGGATGGTG CCCGTAAAAC AAGGACAATN
GGCCTACCAC G	C CCGGATSOTO C
GACTICTITG GO	T CTGAAGAAAC
CCCANANACA GI	T CCGTTTTTGT
781 CCACCTGTAA GGCAAAACA	GGTCGACATT

FIG. 79C

GCCIGGAAT AAATGICATT CGGACCTTTA TITACAGTAA AACCITAAAC CCACGICCAC TIAAATAIG TIGGAATIIG GGIGCAGGIG AAITTIAITAC NNNTACTCCA 841

TCAATCTGTC GTGGAGAATT GANATCAANA ATCTGATATT ATACTGAGAT GITTAGTTAT TAGACTATAA TATGACTCTA CALAICAATA 106

GCAGCATGCT GCTGTGCGGT CACGACCOTC ATGCACTCAG GTGCTGGGAG TACGTGAGTC TCTCTGCGGT AGAGACGCCA CTGTAAGCTT GACATTCGAA 196

CCTGTTTGAG CCCACCAAC AAGCACAGAT TGTTTTCTTC TGCCTGTACA ACAAAAGAAG ACGGACATGT CTGTCATGTC 1021

ACAATCAGTA TGTTAGTCAT CAATAAGGAA GITATICCII ACTGCACATG TGACGTGTAC ATGCANNNN NGATCTTAGA TACGINNNN NCTAGAATCT GAATATGAA CTTTATACTT 1081

ATGCTCTATC TACGAGATAG AATTAACATC TCGTTTTAAA TTAATTGTAG AGCAAATTT AATTCATTAG TTAAGTAA'FC TCTCGTGGAA AGAATCACTT TCTTAGTGAA 1141

1201 AAAGIGIAAA TAATICCICI CICITITICCC ITITICACIA AGGAGITIGI ATAITAAACA	TITCACATIT ATTAACCACA GAGAAAGGG AAAAGTGAT TCCTCAAACA TATAATITGT
AGGACTITGE	TCCTCAAACA
TTTTCACTA	AAAAAGTGAT
CTCTTTTCCC	GAGAAAAGGG
TAATTCCTCT	ATTAAGGAGA
MAGTGTMA	TITCACATIT
1201	

GCCACGTATA	CGCTCCATAT
ACAATAAAAT	TOTTATTTA
TAANNTATT	TTAAAGTTC ATTACATAAT ATTTAAATAA ATTNNATAAA TGTTATTTTA CGGTGCATA!
TAATTTATT	STIAAAGTIC ATTACATAAT ATTIAAATAA A
TAATGTATTA	ATTACATAAT
1261 GAATTTCAAG TAATGTATTA TAAATTTATT TAANNTATTT ACAATAAAAT GCCACGTA	CITANAGITIC
1261	

ANACAGCAG	TTTTGTCGTC
ATACATAGTC	TATOTATCAG
AGAAAGCACA	TCTTTCGTGT
NNNCATTGGT	NNNGTAACCA
AACATGANNN	CG TTGTACTNNN NNNGTAACCA TCTTTCGTGT TATGTATCAG TTTTGTCGTC
1321 AGCATCAAGC AACATGANNN NNNCATTGGT AGAAAGCACA ATACATAGTC AAAACAGCAG	TCGTAGTTCG
1321	

TACTTAATTA	ATGAATTAAT
AATATACATA	TTATATGLAT
1381 AGTATTAAAT AAACAGAAAA TITGCAAAAG GCAAGTAAAG AATATACATA TACTTAATTA	TCATAATITA ITIGICTITI AAACGIIIIC CSIICATITC ITAIAIGIAI AIGAAFIAA:
TITGCARAG	AAACGITITC
AAACAGAAAA	TITCICTITI
AGTATTAAAT	TCATAATTTA
1381	

TGGGGGCAAC	ACCCCCGTTG
AAGCAGATAA	TTCGTCTATT
GANATITAGT	CTTTANATCA
CAGGTAGAAA	CTCCATCTTT
ATTGATACAG	TAACTATGIC
1441 TACATAAAAT ATTGATACAG GAGGTAGAAA GAAATTTAGT AAGCAGATAA TGGGGGCAAC	ATCIATITIA TAACIATGIC CICCATCIII CITIAAAICA IICGICIATI ACCCCCGGIIG
1441	

1561 CTAACAAAA GCAGCCTGAA AAATCGAGCT GCAAACATAG ATTAGCAATC GGCTGAAAGT

FIG. 79E

GATTOTITIT CGTCGGACIT TITAGCICGA CGITIGIAIC TAATCGITAG CCGACITICA

CCGCCCACCG OGCGCGTGGC CCTGGAGCCG GGACCTCGGC GEGCCAATAG TAAAGGGCTA CACGGTTATC ATTTCCCGAT GCTGGCAGCT CGCCCTCTTA GCGGGAGNAT 1621

DAGGTCGGGA CGGATCACCT ATCCCAGCAC TITGGGAGGG CGAGGCAACG TAGGGTCGTG AAACCCTCCC GCTCCGTTGC TCACGCTGTA

1681

1741

AAAAAAAAA TACTAMAMA AGCCCGACCA ACATGGAGAA ACCCCGTCTC TCGGGCTGGT TGTACCTCTT TGGGGCAGAG GTTTGAGATC CAAACTCTAG

129/130

CTGAGGCAGG CACATCCCAG GTGTAGGGTC GCATGGTGGC ACATGCCTTG CGTACCACCG TGTACGGAAC AATGAGCCGG TTACTCGGCC AAAGGCAAAA TTTCCGTTTT

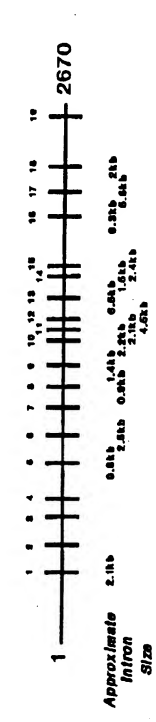
1801

TCATTGCACT CGAGATCACG AGGTAGAGAT TGCGGTGAAG TCCATCTCTA ACGCCACTTC TGAACCTGGG ACTTGGACCC TCTTAAGTGA AGAATTCACT 1961

GAAAAA AAAANNCAAA TTTTNNGTTT CCAGCCTGGG 1921

Genomic Organization of PSM Gene

FIG. 80



### INTERNATIONAL SEARCH REPORT

International application No. PCT/US96/02424

A. CLASSIFICATION OF SUBJECT MATTER  IPC(6) :C12N 15/12, 15/64; C12Q 1/68; C07K 14/435  US CL :536/23.5; 435/6, 7.1, 320.1, 252.3, 69.3; 530/350  According to International Patent Classification (IPC) or to both national classification and IPC			
B. FIELDS SEARCHED			
Minimum documentation searched (classifie	cation system followed by cla	ssification symbols)	
U.S. : 536/23.5; 435/6, 7.1, 320.1, 2	52.3, 69.3; 530/350		
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched			
Electronic data base consulted during the ir	nternational search (name of c	ata base and, where practicable	, search terms used)
INPADOC, CA search terms: prostate specific mem	brane antigen		
C. DOCUMENTS CONSIDERED TO	BE RELEVANT		
Category* Citation of document, with	n indication, where appropriate	e, of the relevant passages	Relevant to claim No.
	WO, A, 94/09820 (SLOAN-KETTERING INSTITUTE FOR CANCER RESEARCH) 11 May 1994, see entire document.		
·			
Further documents are listed in the c	continuation of Box C.	See patent family annex.	
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"P" document published prior to the international	al filling date but later than •&•	document member of the same paten	
Date of the actual completion of the interna	itional search Date o	f mailing of the international se	arch report
29 APRIL 1996 14 MAY 1996			
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Facsimile No. (703) 305-3230	Teleph	one No. (703) 308-0196	